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OM protein - protein search, using sw model

Run on: July 4, 2004, 04:30:15; Search time 39.8358 Seconds

(without alignments)

120.578 Million cell updates/sec

Title: US-09-641-802-2

Perfect score: 89

Sequence: 1 LQTPQPLLQVMMEPQGD 17

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

1586107 seqs, 282547505 residues Searched:

Total number of hits satisfying chosen parameters: 422553

Minimum DB seq length: 7 Maximum DB seq length: 18

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 100 summaries

Database : A Geneseq 29Jan04:\*

1: geneseqp1980s:\*

2: geneseqp1990s:\*

3: geneseqp2000s:\* 4: geneseqp2001s:\*

5: geneseqp2002s:\*

6: geneseqp2003as:\*

7: geneseqp2003bs:\*

8: geneseqp2004s:\*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

## SUMMARIES

		8				
Result		Query				
No.	Score	Match	Length DB		ID	Description
1	 89	100.0	 17	4	AAB72501	Aab72501 Colostrin
2	89	100.0	17	4	AAB59310	Aab59310 Ewe colos
3	89	100.0	17	4	AAB72247	Aab72247 Colostrin
4	89	100.0	17	4	AAB72533	Aab72533 Colostrin
5	89	100.0	17	5	AAO14578	Aao14578 Neural ce
6	89	100.0	17	5	AAM51037	Aam51037 Colostrin
7	89	100.0	17	5	AAE20229	Aae20229 Colostrin
8	89	100.0	18	4	AAB59341	Aab59341 Ewe colos
9	56	62.9	11	4	AAE07185	Aae07185 Colostrin

10	56	62.9	12	4	AAE07195	Aae07	195	Modified
11	34	38.2	15	5	ABB99038	Abb99	038	Carbamyla
12	33	37.1	12	7	ADC44467	Adc44	467	Endotheli
13	33	37.1	15	4	AAB72531	Aab72	:531	Colostrin
14	33	37.1	15	4	AAB59334	Aab59	334	Ewe colos
15	33	37.1	15	4	AAB72279			Colostrin
16	33	37.1	15	4	AAB72563	Aab72	563	Colostrin
17	33	37.1	15	5	AAO14610	Aao14	610	Neural ce
18	33	37.1	15	5	AAM51066			Colostrin
19	33	37.1	15	5	AAE20261			Colostrin
20	32	36.0	8	2	AAR21055			Gamma-car
21	32	36.0	15	6	ABR31148			Human can
22	32	36.0	15	6	ABR31149			Human can
23	32	36.0	15	6	ABR30440			Human can
24	32	36.0	15	6	ABR31517			Human can
25	32	36.0	15	6	ABR30536			Human can
26	32	36.0	15	6	ABR30847			Human can
27	31	34.8	12	3	AAB23189			Hsp47-bin
28	31	34.8	14	4	AAM97279			Human pep
29	31	34.8	15	2	AAW56352			LO-CD2a l
30	31	34.8	15	2	AAY30191			Framework
31	31	34.8	15	5	AAU74511			Human ATP
32	31	34.8	15	6	ABR38577			Human can
33	31	34.8	15	6	ABR38568			Human can
34	31	34.8	15	6	ABR38729			Human can
35	30	33.7	10	7	ADD15523			Predicted
36	30	33.7	14	4	AAM96749			Human pep
37	30	33.7	15	5	AAM47320			Human zin
38	30	33.7	15	6	ABR30169			Human can
39	30	33.7	15	6	ABR30262			Human can
40	30	33.7	15	6	ABR30274			Human can
41	30	33.7	15	6	ABR30229			Human can
42	30	33.7	15	6	ABR30154			Human can
43	30	33.7	15	6	ABR31758			Human can
44	30	33.7	15	6	ABR30297			Human can
45	30	33.7	16	3	AAY65560			Oestrogen
46	30	33.7	16	5	AAU86297			Oestrogen
47	30	33.7	17	4	AAU01845			Wheat pep
48	30	33.7	17	4	AAU01825			Wheat Gli
49	29	32.6	8	6	ABP73085			Amino aci
50	29	32.6	9	5	ABB08361			Synthetic
51	29	32.6	9	7	ABR82213			Human ant
52	29	32.6	10	2	AAY05729			Human G p
53	29	32.6	10	6	ABR05972			Human can
54	29	32.6	10	6	ABR05795			Human can
55	29	32.6	10	6	ABR05791			Human can
56	29	32.6	10	6	ADA14236			Human GPR
57	29	32.6	12	3	AAY92994			Transform
58	29	32.6	12	6	ABU07930			Neisseria
59	29	32.6	13	6	ABP81099			Human TPO
60	29	32.6	15	3	AAY93031	<del>-</del>		Transform
61	29	32.6	15	3	AAY93032	<del>-</del>		Transform
62	29	32.6	15	5	ABB09888			N-termina
63	29	32.6	15	6	ABR31395			Human can
64	29	32.6	15	6	ABR30682			Human can
65	29	32.6	15	6	ABR31783			Human can
66	29	32.6	15	6	ABR31394			Human can
- •		52.0		-				

```
67
        29
             32.6
                     18 2 AAW66658
                                                     Aaw66658 HSV-2 gly
68
        29
             32.6
                     18 4 AAM20225
                                                     Aam20225 Peptide #
69
        29
             32.6
                     18 4 ABB40529
                                                     Abb40529 Peptide #
70
        29
             32.6
                     18 4 AAM34255
                                                     Aam34255 Peptide #
                     18 4 ABB24849
71
        29
             32.6
                                                     Abb24849 Protein #
                     18 4 AAM74093
72
        29
             32.6
                                                     Aam74093 Human bon
73
        29
             32.6
                     18 4 AAM61355
                                                     Aam61355 Human bra
74
        29
             32.6
                     18 4 ABG55849
                                                     Abg55849 Human liv
75
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        29
             32.6
                                                     Abg43991 Human pep
76
             31.5
                     7 2 AAW17515
                                                    Aaw17515 Protein k
        28
77
             31.5
                      9 7 ADD99989
        28
                                                     Add99989 Human 193
                      9 7
78
                                                     Add99556 Human 193
        28
             31.5
                            ADD99556
                      9 7
79
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                                                    Add99875 Human 193
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             31.5
                      9
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                            ADD99363
                                                    Add99363 Human 193
                      9 7 ADD99504
             31.5
                                                     Add99504 Human 193
81
        28
                      9 7 ADD97043
82
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             31.5
                                                     Add97043 Human 193
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                      9 7 ADD98435
                                                     Add98435 Human 193
84
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             31.5
                      9
                         7 ADD99089
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86
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             31.5
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                                                    Add97071 Human 193
87
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             31.5
                         7 ADD98725
88
             31.5
                      9
                        7 ADD98160
                                                   Add98160 Human 193
        28
                      9
                        7 ADD99400
89
             31.5
                                                   Add99400 Human 193
        28
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90
        28
             31.5
                      9 7 ADD99653
91
        28
             31.5
                      9 7 ADD99789
                                                   Add99789 Human 193
                      9 7 ADE00025
                                                    Ade00025 Human 193
92
        28
             31.5
                      9 7
                                                    Add97346 Human 193
93
        28
             31.5
                            ADD97346
                                                   Aag87925 Saccharom
94
        28
             31.5
                     10 4
                            AAG87925
95
        28
             31.5
                     10 7
                                                   Add98298 Human 193
                            ADD98298
96
        28
             31.5
                     10 7
                                                    Ade00501 Human 193
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97
        28
             31.5
                     10 7
                            ADD98062
                                                    Add98062 Human 193
98
             31.5
                     10 7 ADD98902
                                                    Add98902 Human 193
        28
                     10 7 ADD97476
99
             31.5
                                                    Add97476 Human 193
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                     10 7 ADE00482
        28
             31.5
                                                     Ade00482 Human 193
100
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## ALIGNMENTS

```
RESULT 1
AAB72501
     AAB72501 standard; peptide; 17 AA.
ΙD
XX
AC
    AAB72501;
XX
     09-MAY-2001 (first entry)
DT
XX
DE
     Colostrinin peptide #2.
XX
KW
     Dermatological; oxidative stress regulator; colostrinin.
XX
     Unidentified.
OS
XX
PN
     W0200112650-A2.
XX
PD
     22-FEB-2001.
XX
```

```
PF
    17-AUG-2000; 2000WO-US022665.
XX
PR
    17-AUG-1999;
                   99US-0149310P.
XX
PΑ
     (TEXA ) UNIV TEXAS SYSTEM.
XX
PΙ
    Stanton GJ, Hughes TK, Boldogh I;
XX
DR
    WPI; 2001-218342/22.
XX
PT
    Modulating oxidative stress level in a cell, involves contacting the cell
PT
    with an oxidative stress regulator selected from colostrinin, its
PT
    constituent peptide, analog or their combinations.
XX
PS
    Claim 6; Page 25; 48pp; English.
XX
CC
    The present invention relates to a method for modulating the oxidative
CC
    stress level in a cell or a patient, comprising contacting the cell with,
    or administering to the patient, an oxidative stress regulator selected
CC
CC
    from colostrinin, or its constituent peptide (e.g. the present peptide),
CC
    to change the level of an oxidising species in the cell. The method can
CC
    be used to treat oxidative damage to skin, by decreasing or preventing an
CC
    increase in the level of damage to a biomolecule of the patient
XX
SQ
    Sequence 17 AA;
 Query Match
                         100.0%; Score 89; DB 4; Length 17;
 Best Local Similarity 100.0%; Pred. No. 2.4e-07;
                              0; Mismatches 0; Indels
 Matches
           17; Conservative
                                                                            0;
                                                                0; Gaps
           1 LOTPOPLLOVMMEPOGD 17
Qy
             1 LQTPQPLLQVMMEPQGD 17
Db
RESULT 2
AAB59310
ID
    AAB59310 standard; peptide; 17 AA.
XX
AC
    AAB59310;
XX
DT
    21-MAR-2001 (first entry)
XX
DE
    Ewe colostrinin peptide fragment A-1.
XX
KW
     Sheep; colostrinin; proline rich polypeptide; colostrum; immune disorder;
KW
    central nervous system disorder; dietary supplement; beta-amyloid plaque.
XX
os
    Ovis sp.
XX
PN
    WO200075173-A2.
XX
PD
    14-DEC-2000.
XX
PF
     02-JUN-2000; 2000WO-GB002128.
XX
PR
     02-JUN-1999; 99GB-00012852.
```

```
XX
PA
     (REGE-) REGEN THERAPEUTICS PLC.
XX
PΙ
    Georgiades JA;
XX
DR
    WPI; 2001-071058/08.
XX
PΤ
     Peptides having an N-terminal amino acid sequence isolated from
PT
     colostrinin for treating e.g. disorders of the central nervous system and
PT
     immune system, viral and bacterial infections, and diseases characterized
PT
    by amyloid plaques.
XX
    Claim 7; Page 27; 63pp; English.
PS
XX
CC
    The present invention provides the sequences of a number of peptides
CC
     found in ewe's colostrinin. Colostrinin is the proline-rich polypeptide
CC
     fragment of colostrum. These peptides can be used in the treatment of
CC
     central nervous system disorders such as senile dementia, Parkinson's
CC
    disease, Alzheimer's disease, psychosis and neurosis, immune system
CC
    disorders such as bacterial and viral infections, to improve the
    development of a child's immune system, as a dietary supplement, and to
CC
CC
    promote the dissolution of beta-amyloid plaques
XX
SO
     Sequence 17 AA;
 Query Match 100.0%; Score 89; DB 4; Length 17; Best Local Similarity 100.0%; Pred. No. 2.4e-07;
  Matches 17; Conservative
                               0; Mismatches
                                                   0; Indels
                                                                  0; Gaps
                                                                              0;
            1 LQTPQPLLQVMMEPQGD 17
Qу
             Db
            1 LQTPQPLLQVMMEPQGD 17
RESULT 3
AAB72247
    AAB72247 standard; peptide; 17 AA.
XX
AC
    AAB72247;
XX
DΤ
     14-MAY-2001 (first entry)
XX
DE
     Colostrinin derived cytokine inducing peptide SEQ ID 2.
XX
KW
     Colostrinin; immune response; cytokine; blood cell proliferation;
KW
     central nervous system disorder; neurological diosrder; mental disorder;
     dementia; neurodegenerative disease; Alzheimer's disease; psychosis;
KW
KW
     neurosis; infection.
XX
os
     Synthetic.
XX
     WO200111937-A2.
PN
XX
PD
     22-FEB-2001.
XX
PF
     17-AUG-2000; 2000WO-US022818.
XX
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PR
     17-AUG-1999;
                    99US-0149311P.
XX
PA
     (TEXA ) UNIV TEXAS SYSTEM.
PΑ
     (REGE-) REGEN THERAPEUTICS PLC.
XX
PΙ
     Stanton GJ, Hughes TK, Boldogh I, Georgiades J;
XX
    WPI; 2001-202804/20.
DR
XX
PT
    Inducing a cytokine and modulating an immune response, useful for
PT
     treating central nervous system diseases and bacterial and viral
     infections, comprises administering colostrinin as an immunological
PT
PT
     regulator.
XX
PS
    Claim 1; Page 34; 50pp; English.
XX
CC
     Sequences AAB72246 - AAB72275 represent peptides derived from clostrinin,
CC
    a proline rich polypeptide aggregate contained in colostrum. The peptides
    have immune response modulatory activity, and are capable of inducing
CC
CC
    cytokines. Colostrinin and its derived peptides are useful for inducing
CC
    cytokine production, for modulating an immunological response and for
CC
     inducing blood cell proliferation. The peptides are useful in the
CC
     treatment of disorders of the central nervous system, neurological
CC
    disorders, mental disorders, dementia, neurodegenerative diseases,
CC
    Alzheimer's disease, motor neurone disease, psychosis, neurosis, chronic
CC
    disorders of the immune system, bacterial and viral infections and
CC
     acquired immunological deficiencies
XX
SO
     Sequence 17 AA;
  Query Match
                          100.0%; Score 89; DB 4; Length 17;
                         100.0%; Pred. No. 2.4e-07;
  Best Local Similarity
 Matches
           17; Conservative
                              0; Mismatches
                                                  0; Indels
                                                                             0;
                                                                 0; Gaps
Qу
           1 LOTPOPLLOVMMEPOGD 17
              11111111111111111
Dh
           1 LQTPQPLLQVMMEPQGD 17
RESULT 4
AAB72533
    AAB72533 standard; peptide; 17 AA.
ΙD
XX
AC
    AAB72533;
XX
DT
     09-MAY-2001 (first entry)
XX
DE
    Colostrinin peptide #2.
XX
KW
     Neuroprotective; neural cell differentiation regulator; colostrinin;
KW
     colostrum.
XX
os
     Unidentified.
XX
PN
    WO200112651-A2.
XX
PD
     22-FEB-2001.
```

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XX
PF
     17-AUG-2000; 2000WO-US022774.
XX
     17-AUG-1999;
                    99US-0149633P.
PR
XX
PA
     (TEXA ) UNIV TEXAS SYSTEM.
XX
PΙ
     Boldogh I;
XX
DR
     WPI; 2001-226545/23.
XX
PT
     Use of colostrinin, its constituent peptide or analog as a neural cell
PT
     regulator, for promoting neural cell differentiation and treating damaged
PT
     neural cells in a patient.
XX
PS
     Claim 6; Page 21; 35pp; English.
XX
CC
     The present invention relates to a method for promoting neural cell
CC
     differentiation and treating damaged neural cells, using colostrinin and
CC
     colostrinin constituent peptides (e.g. the present peptide) as a neural
CC
     cell regulator. Colostrinin is a polypeptide complex found in colostrum
XX
SQ
     Sequence 17 AA;
  Query Match
                          100.0%; Score 89; DB 4; Length 17;
  Best Local Similarity 100.0%; Pred. No. 2.4e-07;
  Matches
            17; Conservative
                               0; Mismatches 0; Indels
                                                                 0; Gaps
                                                                              0;
            1 LQTPQPLLQVMMEPQGD 17
Qу
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Db
            1 LQTPQPLLQVMMEPQGD 17
RESULT 5
AAO14578
     AAO14578 standard; peptide; 17 AA.
XX
AC
    AAO14578;
XX
     27-MAY-2002 (first entry)
DT
XX
DE
     Neural cell regulatory colostrinin peptide 2.
XX
KW
     Neural cell differentiation; neural cell regulator; colostrinin peptide;
KW
     neural cell formation; proline-rich polypeptide aggregate; colostrum;
KW
     neural cell treatment.
XX
os
     Unidentified.
XX
FH
                     Location/Qualifiers
     Key
FT
     Modified-site
                     /note= "Optional C-terminal amide"
FT
XX
PN
     WO200213851-A1.
XX
PD
     21-FEB-2002.
XX
```

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ΡF
     17-AUG-2000; 2000WO-US022777.
XX
PR
     17-AUG-2000; 2000WO-US022777.
XX
     (TEXA ) UNIV TEXAS SYSTEM.
PA
XX
     Boldogh I, Stanton JG, Hughes TK;
PΙ
XX
DR
    WPI: 2002-269152/31.
XX
     Promoting cell differentiation in a patient involves use of blood cell
PΤ
     regulator selected from colostrinin, its constituent peptide and/or
PT
PT
     analog.
XX
ΡŞ
     Claim 7; Page 21; 37pp; English.
XX
     The invention comprises a method for promoting cell differentiation (e.g.
CC
     neural cell differentiation). The method involves contacting cells with a
CC
     neural cell regulator (i.e. a colostrinin peptide) in order to change the
CC
     cells in morphology to form neural cells. Colostrinin is a proline-rich
CC
     polypeptide aggregate that is present in colostrum. The method of the
CC
     invention is useful for promoting the differentiation of cells and for
CC
CC
     treating damaged neural cells in a patient. The present amino acid
     sequence represents a specifically claimed colostrinin peptide used in
CC
CC
     the method of the invention
XX
SO
     Sequence 17 AA;
                          100.0%; Score 89; DB 5; Length 17;
  Query Match
                          100.0%; Pred. No. 2.4e-07;
  Best Local Similarity
           17; Conservative
                                0; Mismatches
                                                 0; Indels
                                                                 0; Gaps
                                                                             0;
  Matches
            1 LQTPQPLLQVMMEPQGD 17
Qу
              Db
            1 LQTPQPLLQVMMEPQGD 17
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AAM51037
     AAM51037 standard; peptide; 17 AA.
ID
XX
AC
     AAM51037;
XX
DT
     30-MAY-2002 (first entry)
XX
DE
     Colostrinin constituent peptide.
XX
     Colostrinin; colostrum; immunomodulator; cardiovascular;
KW
     blood cell regulator; cytokine inducer; human.
KW
XX
OS
     Homo sapiens.
XX
FH
                     Location/Qualifiers
FT
     Modified-site
                     17
FT
                     /note= "optional C-terminal amidation"
XX
PN
     W0200213849-A1.
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XX
PD
     21-FEB-2002.
XX
PF
     17-AUG-2000; 2000WO-US022775.
XX
PR
     17-AUG-2000; 2000WO-US022775.
XX
     (TEXA ) UNIV TEXAS SYSTEM.
PA
PA
     (REGE-) REGEN THERAPEUTICS PLC.
XX
     Stanton GJ, Hughes TK, Boldogh I, Georgiades J;
PΙ
XX
DR
     WPI; 2002-269150/31.
XX
PT
     Modulation of blood cell proliferation in a patient involves use of blood
PT
     cell regulator selected from colostrinin, its constituent peptide and/or
PТ
     analog.
XX
PS
     Claim 1; Page 34; 54pp; English.
XX
CC
     The present sequence is that of a colostrinin constituent peptide that is
CC
     preferred for use as an immunological regulator and as a blood cell
     regulator in claimed methods of the invention. Methods are claimed for:
CC
CC
     inducing a cytokine in a cell by contact with an immunological regulator,
CC
     where the cell is present in a cell culture, a tissue, an organ or an
CC
     organism, and the cell is mammalian, including human; modulating an
     immune response in a cell by contact with the immunological regulator
CC
CC
     under conditions effective to induce a cytokine; modulating an immune
CC
     response in a patient by administering an immunological regulator under
     conditions effective to induce a cytokine, where the immunological
CC
CC
     regulator is administered topically or as part of a dietary supplement,
     and where the immune response is specific or non specific, an interferon
CC
CC
     response or an antibody response; modulating blood cell proliferation by
     contacting blood cells with a blood cell regulator, where the blood cells
CC
CC
     are present in a cell culture or an organism, are mammalian or human, and
     where the blood cells are increased in number or differentiated; and a
CC
CC
     method for modulating blood cell proliferation in a patent. A claimed
CC
     cytokine-inducing composition comprises a pharmaceutical carrier and an
CC
     active agent such as the present peptide. Cytokines induced by this
CC
     peptide in human leucocyte cultures include interferon-gamma, tumour
CC
     necrosis factor-alpha, interleukin-4, interleukin-6, interleukin-10 and
CC
     interleukin-12. It was one of the best overall inducers in almost all
CC
     cytokine and blood cell proliferation experiments conducted
XX
SQ
     Sequence 17 AA;
  Query Match
                          100.0%; Score 89; DB 5; Length 17;
                          100.0%; Pred. No. 2.4e-07;
  Best Local Similarity
                                0; Mismatches
                                                                             0;
  Matches
           17; Conservative
                                                  0; Indels
                                                                 0; Gaps
            1 LOTPOPLLOVMMEPQGD 17
Qу
              111111111111111
Db
            1 LQTPQPLLQVMMEPQGD 17
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ID
     AAE20229 standard; peptide; 17 AA.
XX
AC
    AAE20229;
XX
DT
     18-JUN-2002
                 (first entry)
XX
DE
     Colostrinin constituent peptide #2.
XX
KW
     Blood cell regulator; colostrinin; constituent peptide; oxidative stress;
KW
     therapy; oxidative damage; skin; aging; wound healing; cell replacement;
KW
     tissue; organ; cosmetic procedure; repair; regeneration; preservation;
KW
     transplantation; implantation; dermatological; vulnerary.
XX
os
     Unidentified.
XX
FH
                     Location/Qualifiers
     Key
    Modified-site
FT
                     /note= "Optionally C-terminal amide"
FT
XX
     WO200213850-A1.
PN
XX
PD
     21-FEB-2002.
XX
PF
     17-AUG-2000; 2000WO-US022776.
XX
PR
     17-AUG-2000; 2000WO-US022776.
XX
PA
     (TEXA ) UNIV TEXAS SYSTEM.
XX
PΙ
     Stanton GJ, Hughes TK,
                              Boldogh I;
XX
DR
     WPI; 2002-269151/31.
XX
PT
     Composition useful for the modulation of blood cell proliferation in a
PT
     patient comprises a blood cell regulator selected from colostrinin, its
PT
     constituent peptide and/or analog.
XX
PS
     Claim 6; Page 25; 51pp; English.
XX
CC
     The invention relates to a composition which comprises a blood cell
CC
     regulator selected from colostrinin, its constituent peptide and/or
CC
     analogue. The invention is used for modulating the oxidative stress level
CC
     in a cell e.g. mammalian or human cell present in a cell culture, tissue,
CC
     organ, or organism; or for treating oxidative damage to the skin of a
CC
     patient e.g. animal or human; to modulate oxidative stress during/ after
CC
     a premature birth or normal birth, preventing/delaying aging in a
CC
     patient, enhancing wound healing, and the reduction of side effects of
CC
     cosmetic procedures. The method changes the level of an oxidising species
CC
     in the cell, such as decreases or prevents increase in the level of
CC
     damage to a biomolecule of the patient selected from DNA, protein and/or
CC
     lipid, compared to the same conditions when the oxidative stress
CC
     regulator is not present. The modulation of oxidative stress results in
CC
     enhanced repair, regeneration, and replacement of cells, tissues and
CC
     organs (e.g. kidney, liver, pancreas, skin, and the other internal and
CC
     external organs), as well as enhanced preservation of such organs for
CC
     transplantation, implantation, or scientific research. The present
```

sequence is a colostrinin constituent peptide

CC

```
XX
SQ
     Sequence 17 AA;
  Query Match
                          100.0%; Score 89; DB 5; Length 17;
                          100.0%; Pred. No. 2.4e-07;
  Best Local Similarity
                                0; Mismatches
            17; Conservative
                                                   0; Indels
                                                                  0; Gaps
                                                                              0;
            1 LQTPQPLLQVMMEPQGD 17
Qy
              111111111111111
Db
            1 LQTPQPLLQVMMEPQGD 17
RESULT 8
AAB59341
    AAB59341 standard; peptide; 18 AA.
XX
AC
    AAB59341;
XX
DT
     21-MAR-2001 (first entry)
XX
DE
     Ewe colostrinin peptide fragment derived sequence #1.
XX
KW
     Sheep; colostrinin; proline rich polypeptide; colostrum; immune disorder;
KW
     central nervous system disorder; dietary supplement; beta-amyloid plaque.
XX
os
     Ovis sp.
XX
     WO200075173-A2.
PN
XX
     14-DEC-2000.
PD
XX
     02-JUN-2000; 2000WO-GB002128.
PF
XX
     02-JUN-1999;
PR
                    99GB-00012852.
XX
PA
     (REGE-) REGEN THERAPEUTICS PLC.
XX
PΙ
     Georgiades JA;
XX
DR
     WPI; 2001-071058/08.
XX
PT
     Peptides having an N-terminal amino acid sequence isolated from
PT
     colostrinin for treating e.g. disorders of the central nervous system and
PT
     immune system, viral and bacterial infections, and diseases characterized
PT
     by amyloid plaques.
XX
PS
     Claim 8; Page 27; 63pp; English.
XX
CC
     The present invention provides the sequences of a number of peptides
CC
     found in ewe's colostrinin. Colostrinin is the proline-rich polypeptide
     fragment of colostrum. These peptides can be used in the treatment of
CC
     central nervous system disorders such as senile dementia, Parkinson's
CC
CC
     disease, Alzheimer's disease, psychosis and neurosis, immune system
CC
     disorders such as bacterial and viral infections, to improve the
CC
     development of a child's immune system, as a dietary supplement, and to
CC
     promote the dissolution of beta-amyloid plaques
```

XX

```
SO
     Sequence 18 AA;
  Query Match
                          100.0%; Score 89; DB 4; Length 18;
  Best Local Similarity
                          100.0%; Pred. No. 2.6e-07;
                                0; Mismatches
  Matches
           17; Conservative
                                                  0; Indels
                                                                 0; Gaps
            1 LQTPQPLLQVMMEPQGD 17
Qу
              11111111111111
Db
            2 LQTPQPLLQVMMEPQGD 18
RESULT 9
AAE07185
ID
     AAE07185 standard; peptide; 11 AA.
XX
AC
     AAE07185;
XX
     06-NOV-2001 (first entry)
DT
XX
DE
    Colostrinin peptide 1.
XX
KW
     Colostrinin; nootropic; neuroprotective; immunomodulatory; antibacterial;
KW
     Parkinson's disease; Alzheimer's disease; mental disorder; food additive;
KW
     central nervous system disorder; neurodegenerative disorder; weight loss;
KW
     beta-amyloid plaque; psychosis; neurosis; cachexia; motor neuron disease;
KW
     acquired immunological deficiency; neurological disorder; dementia;
KW
     antiviral.
XX
OS
     Unidentified.
XX
     WO200155199-A1.
PN
XX
     02-AUG-2001.
PD
XX
PF
     26-JAN-2001; 2001WO-GB000329.
XX
     26-JAN-2000; 2000GB-00001825.
PR
XX
PA
     (REGE-) REGEN THERAPEUTICS PLC.
XX
PΙ
     Georgiades JA;
XX
DR
     WPI; 2001-488775/53.
XX
PT
     Peptide useful as an interalia in the treatment of e.g. disorders of the
PТ
     immune system and the central nervous system comprises ten amino-terminal
PT
     amino acid sequence derived from peptides present in colostrinin.
XX
PS
     Claim 1; Page 15; 40pp; English.
XX
CC
     The invention relates to colostrinin peptide fragments which are useful,
CC
     inter alia, in the treatment of chronic disorders of the immune system
CC
     and the central nervous system. Colostrinin peptides are used as a
CC
     medicament in the treatment of neurological disorders e.g., dementia,
CC
     neurodegenerative disorders e.g., Alzheimer's disease, motor neuron
CC
     disease e.g., Parkinson's disease, mental disorders e.g. psychosis and
```

neurosis, in acquired immunological deficiencies, chronic bacterial and

CC

```
CC
     viral infections and diseases characterised by the presence of beta-
CC
     amyloid plaques and as a dietary supplement for babies, small children,
    adults and senile persons, who have been subjected to chemotherapy or
CC
    have suffered from cachexia or weight loss due to the chronic disease.
CC
    Colostrinin peptides are also used as food additives and as an auxillary
CC
    withdrawal treatment for drug addicts, after a period of detoxification
CC
    and in persons dependent on stimulants. Colostrinin peptides are used to
CC
    prepare antibodies and to treat emotional disturbances, e.g. emotional
CC
    disturbances of psychiatric patients in a state of depression. These
CC
CC
     colostrinin peptides improves the development of immune system in a new
    born child and to correct the immunological deficiencies in a child. The
CC
CC
    present sequence is colostrinin peptide 1 related to the invention
XX
SQ
     Sequence 11 AA;
  Query Match
                          62.9%; Score 56; DB 4; Length 11;
 Best Local Similarity
                         100.0%; Pred. No. 0.03;
                              0; Mismatches
                                                  0; Indels
                                                                 0; Gaps
 Matches
           11; Conservative
                                                                             0;
            5 QPLLQVMMEPQ 15
Qу
             Db
           1 QPLLQVMMEPQ 11
RESULT 10
AAE07195
    AAE07195 standard; peptide; 12 AA.
ID
XX
AC
    AAE07195;
XX
     06-NOV-2001 (first entry)
DT
XX
DE
    Modified colostrinin cyclic peptide #1.
XX
ΚW
    Colostrinin; nootropic; neuroprotective; immunomodulatory; antibacterial;
KW
     Parkinson's disease; Alzheimer's disease; mental disorder; food additive;
KW
     central nervous system disorder; neurodegenerative disorder; weight loss;
KW
     beta-amyloid plaque; psychosis; neurosis; cachexia; motor neuron disease;
KW
     acquired immunological deficiency; neurological disorder; dementia;
     antiviral; cyclic.
KW
XX
OS
     Synthetic.
XX
FH
                     Location/Qualifiers
     Key
FT
     Modified-site
FT
                     /note= "N-terminal acetyl; this residue forms a cyclic
FT
                     linkage with Gln found at the C-terminal end"
XX
PN
     WO200155199-A1.
XX
PD
     02-AUG-2001.
XX
PF
     26-JAN-2001; 2001WO-GB000329.
XX
PR
     26-JAN-2000; 2000GB-00001825.
XX
PA
     (REGE-) REGEN THERAPEUTICS PLC.
```

```
XX
PΙ
     Georgiades JA;
XX
DR
     WPI; 2001-488775/53.
XX
PT
     Peptide useful as an interalia in the treatment of e.g. disorders of the
     immune system and the central nervous system comprises ten amino-terminal
PT
PT
     amino acid sequence derived from peptides present in colostrinin.
XX
PS
     Example 2; Page 8; 40pp; English.
XX
CC
     The invention relates to colostrinin peptide fragments which are useful,
CC
     inter alia, in the treatment of chronic disorders of the immune system
CC
     and the central nervous system. Colostrinin peptides are used as a
CC
     medicament in the treatment of neurological disorders e.g., dementia,
     neurodegenerative disorders e.g., Alzheimer's disease, motor neuron
CC
CC
     disease e.g., Parkinson's disease, mental disorders e.g. psychosis and
CC
     neurosis, in acquired immunological deficiencies, chronic bacterial and
CC
     viral infections and diseases characterised by the presence of beta-
CC
     amyloid plaques and as a dietary supplement for babies, small children,
CC
     adults and senile persons, who have been subjected to chemotherapy or
CC :
     have suffered from cachexia or weight loss due to the chronic disease.
CC
     Colostrinin peptides are also used as food additives and as an auxillary
CC
     withdrawal treatment for drug addicts, after a period of detoxification
     and in persons dependent on stimulants. Colostrinin peptides are used to
CC
CC
     prepare antibodies and to treat emotional disturbances, e.g. emotional
CC
     disturbances of psychiatric patients in a state of depression. These
CC
     colostrinin peptides improves the development of immune system in a new
CC
     born child and to correct the immunological deficiencies in a child. The
     present sequence is modified colostrinin cyclic peptide #1 related to the
CC
CC
     invention
XX
SO
     Sequence 12 AA;
  Query Match
                          62.9%; Score 56; DB 4; Length 12;
  Best Local Similarity
                          100.0%; Pred. No. 0.032;
  Matches
           11; Conservative
                                0; Mismatches
                                                   0; Indels
                                                                             0;
                                                                 0; Gaps
Qу
            5 QPLLQVMMEPQ 15
              Db
            2 QPLLQVMMEPQ 12
RESULT 11
ABB99038
ID
     ABB99038 standard; peptide; 15 AA.
XX
AC
     ABB99038;
XX
DT
     24-JAN-2003 (first entry)
XX
DΕ
     Carbamylaspartic dehydrase 9.68 N-terminal amino acid sequence.
XX
     Carbamylaspartic dehydrase 9.68; enzyme; malignant tumour; haemopathy;
KW
KW
     development disorder; inflammation; human immunodeficiency virus; HIV;
     infection; immunological disease.
KW
XX
```

```
os
     Unidentified.
XX
PN
     CN1352302-A.
XX
PD
     05-JUN-2002.
XX
PF
     02-NOV-2000; 2000CN-00127191.
XX
PR
     02-NOV-2000; 2000CN-00127191.
XX
     (BODE-) BODE GENE DEV CO LTD SHANGHAI.
PA
XX
     Mao Y, Xie Y;
PΙ
XX
DR
     WPI; 2002-644474/70.
XX
PT
     New polypeptide-carbamylaspartic dehydrase 9.68 and polynucleotide
PT
     encoding the polypeptide.
XX
PS
     Example 5; Page 18 (disclosure); 31pp; Chinese.
XX
CC
     The invention relates to the polypeptide carbamylaspartic dehydrase 9.68,
     the polynucleotides encoding this polypeptide, and a DNA recombination
CC
CC
     process for production of the polypeptide. Also disclosed is the method
CC
     of applying the polypeptide in the treatment of various diseases such as,
     malignant tumours, haemopathy, development disorder, inflammations, human
CC
     immunodeficiency virus (HIV) infection, and immunological diseases. The
CC
     present invention also discloses the antagonist resisting the polypeptide
CC
     and its treatment effect, and the application of the polynucleotides for
CC
     encoding carbamylaspartic dehydrase 9.68. The current sequence represents
CC
CC
     the carbamylaspartic dehydrase 9.68 N-terminal amino acid sequence
XX
SQ
     Sequence 15 AA;
  Query Match
                          38.2%; Score 34; DB 5; Length 15;
                          41.7%; Pred. No. 1.4e+02;
  Best Local Similarity
                                                 3; Indels
  Matches
             5; Conservative
                                 4; Mismatches
                                                                 0; Gaps
                                                                              0;
            3 TPQPLLQVMMEP 14
Qу
                    :::
              : | | | |
Db
            4 SPQPLTGALLQP 15
RESULT 12
ADC44467
     ADC44467 standard; peptide; 12 AA.
XX
АÇ
     ADC44467;
XX
DT
     18-DEC-2003 (first entry)
XX
DE
     Endothelial cell binding peptide SEQ ID NO:195.
XX
     endothelial cell binding protein; ECBP; anti-tumour; cytostatic;
KW
KW
     vasotropic; antipsoriatic; dermatological; ophthalmological;
     antidiabetic; antiarthritic; vulnerary; antiulcer; antiinflammatory;
KW
KW
     antibacterial; gynaecological; angiogenesis.
```

```
XX
OS
     Synthetic.
XX
PN
    WO2003037172-A2.
XX
     08-MAY-2003.
PD
XX
     01-NOV-2002; 2002WO-US035258.
PF
XX
PR
     01-NOV-2001; 2001US-0334822P.
XX
PA
     (GPCB-) GPC BIOTECH INC.
XX
PΙ
     Gyuris J, Lamphere L, Morris AJ, Tsaioun K;
XX
    WPI; 2003-482072/45.
DR
XX
PT
     Novel synthetic or recombinant polypeptide useful for promoting, reducing
     proliferation and/or migration of endothelial cells, and for modulating
PT
     angiogenesis, has endothelial cell binding protein sequences.
PT
XX
PS
     Claim 3; SEQ ID NO 195; 126pp; English.
XX
CC
     The invention relates to a novel isolated, synthetic or recombinant
CC
     peptide or polypeptide which includes one or more endothelial cell
CC
     binding protein (ECBP) sequences. A peptide of the invention has anti-
CC
     tumour, cytostatic, vasotropic, antipsoriatic, dermatological,
CC
     ophthalmological, antidiabetic, antiarthritic, vulnerary, antiulcer,
CC
     antiinflammatory, antibacterial, and gynaecological activity. The peptide
     is useful for promoting, reducing the proliferation and/or migration of
CC
CC
     endothelial cells, by treating the cells with an ECBP agonist, which is
CC
     preferably the peptide, to promote proliferation and/or migration of the
CC
     treated cells, and for reducing or promoting angiogenesis, by treating
     the cells with an ECBP antagonist, which is preferably the peptide of the
CC
CC
     invention. A peptide of the invention is also useful for manufacturing a
CC
     medicament for promoting angiogenesis, by admixing an ECBP agonist or
CC
     ECBP antagonist to promote or reduce angiogenesis at one or more sites in
     a treated mammal. The medicament is useful for promoting or reducing
CC
     angiogenesis. ECBP sequences are useful to alter the infectivity spectrum
CC
CC
     of a viral particle. The present sequence represents an ECBP of the
CC
     invention.
XX
SO
     Sequence 12 AA;
                          37.1%; Score 33; DB 7; Length 12;
  Query Match
  Best Local Similarity
                          100.0%; Pred. No. 1.6e+02;
  Matches
             6; Conservative
                                0; Mismatches
                                                  0; Indels
                                                                 0; Gaps
                                                                              0;
            1 LQTPQP 6
Qу
              +11111
Db
            2 LQTPQP 7
RESULT 13
AAB72531
ΙD
     AAB72531 standard; peptide; 15 AA.
XX
```

```
AAB72531;
AC
XX
DT
     09-MAY-2001 (first entry)
XX
DE
    Colostrinin peptide #32.
XX
KW
     Dermatological; oxidative stress regulator; colostrinin.
XX
os
    Unidentified.
XX
PN
    WO200112650-A2.
XX
PD
     22-FEB-2001.
XX
    17-AUG-2000; 2000WO-US022665.
PF
XX
PR
    17-AUG-1999;
                    99US-0149310P.
XX
     (TEXA ) UNIV TEXAS SYSTEM.
PA
XX
PΙ
     Stanton GJ, Hughes TK, Boldogh I;
XX
DR
    WPI; 2001-218342/22.
XX
PT
    Modulating oxidative stress level in a cell, involves contacting the cell
     with an oxidative stress regulator selected from colostrinin, its
PT
PΤ
     constituent peptide, analog or their combinations.
XX
PS
     Claim 6; Page 26; 48pp; English.
XX
    The present invention relates to a method for modulating the oxidative
CC
CC
     stress level in a cell or a patient, comprising contacting the cell with,
CC
     or administering to the patient, an oxidative stress regulator selected
CC
     from colostrinin, or its constituent peptide (e.g. the present peptide),
     to change the level of an oxidising species in the cell. The method can
CC
CC
     be used to treat oxidative damage to skin, by decreasing or preventing an
CC
     increase in the level of damage to a biomolecule of the patient
XX
SO
     Sequence 15 AA;
  Query Match
                          37.1%; Score 33; DB 4; Length 15;
                          53.8%; Pred. No. 2e+02;
  Best Local Similarity
             7; Conservative
                                 1; Mismatches
  Matches
                                                  5; Indels
                                                                 0; Gaps
                                                                             0;
           2 QTPQPLLQVMMEP 14
Qу
             1 1111
                      : | |
Db
            3 QPPQPLPPTVMFP 15
RESULT 14
AAB59334
ID
     AAB59334 standard; peptide; 15 AA.
XX
AC
    AAB59334;
XX
DT
     21-MAR-2001 (first entry)
XX
```

```
DΕ
     Ewe colostrinin peptide fragment C-9.
XX
KW
     Sheep; colostrinin; proline rich polypeptide; colostrum; immune disorder;
KW
     central nervous system disorder; dietary supplement; beta-amyloid plaque.
XX
os
    Ovis sp.
XX
    WO200075173-A2.
PN
XX
PD
     14-DEC-2000.
XX
     02-JUN-2000; 2000WO-GB002128.
PF
XX
PR
     02-JUN-1999;
                    99GB-00012852.
XX
     (REGE-) REGEN THERAPEUTICS PLC.
PA
XX
ΡI
    Georgiades JA;
XX
    WPI; 2001-071058/08.
DR
XX
PT
     Peptides having an N-terminal amino acid sequence isolated from
PT
     colostrinin for treating e.g. disorders of the central nervous system and
PT
     immune system, viral and bacterial infections, and diseases characterized
PT
     by amyloid plaques.
XX
PS
     Claim 7; Page 27; 63pp; English.
XX
CC
     The present invention provides the sequences of a number of peptides
     found in ewe's colostrinin. Colostrinin is the proline-rich polypeptide
CC
CC
     fragment of colostrum. These peptides can be used in the treatment of
CC
     central nervous system disorders such as senile dementia, Parkinson's
CC
     disease, Alzheimer's disease, psychosis and neurosis, immune system
     disorders such as bacterial and viral infections, to improve the
CC
CC
     development of a child's immune system, as a dietary supplement, and to
CC
     promote the dissolution of beta-amyloid plaques
XX
     Sequence 15 AA;
SQ
  Query Match
                          37.1%; Score 33; DB 4; Length 15;
                          53.8%; Pred. No. 2e+02;
  Best Local Similarity
                                 1; Mismatches 5; Indels
  Matches
             7; Conservative
                                                                 0; Gaps
                                                                             0;
            2 QTPQPLLQVMMEP 14
Qу
              1 1111 :1 1
Db
            3 QPPQPLPPTVMFP 15
RESULT 15
AAB72279
ID
     AAB72279 standard; peptide; 15 AA.
XX
AC
     AAB72279;
XX
DT
     14-MAY-2001 (first entry)
XX
DE
     Colostrinin derived cytokine inducing peptide SEQ ID 34.
```

```
XX
KW
     Colostrinin; immune response; cytokine; blood cell proliferation;
KW
     central nervous system disorder; neurological diosrder; mental disorder;
KW
     dementia; neurodegenerative disease; Alzheimer's disease; psychosis;
     neurosis; infection.
KW
XX
OS
     Synthetic.
XX
PN
    WO200111937-A2.
XX
PD
     22-FEB-2001.
XX
     17-AUG-2000; 2000WO-US022818.
PF
XX
PR
     17-AUG-1999;
                    99US-0149311P.
XX
     (TEXA ) UNIV TEXAS SYSTEM.
PA
     (REGE-) REGEN THERAPEUTICS PLC.
PA
XX
PΙ
     Stanton GJ, Hughes TK, Boldogh I, Georgiades J;
XX
DR
    WPI; 2001-202804/20.
XX
PT
     Inducing a cytokine and modulating an immune response, useful for
PT
     treating central nervous system diseases and bacterial and viral
PT
     infections, comprises administering colostrinin as an immunological
PT
     regulator.
XX
PS
     Claim 1; Page 34; 50pp; English.
XX
CC
     Sequences AAB72246 - AAB72275 represent peptides derived from clostrinin,
     a proline rich polypeptide aggregate contained in colostrum. The peptides
CC
     have immune response modulatory activity, and are capable of inducing
CC
     cytokines. Colostrinin and its derived peptides are useful for inducing
CC
CC
     cytokine production, for modulating an immunological response and for
CC
     inducing blood cell proliferation. The peptides are useful in the
     treatment of disorders of the central nervous system, neurological
CC
CC
     disorders, mental disorders, dementia, neurodegenerative diseases,
     Alzheimer's disease, motor neurone disease, psychosis, neurosis, chronic
CC
CC
     disorders of the immune system, bacterial and viral infections and
CC
     acquired immunological deficiencies
XX
SO
     Sequence 15 AA;
                          37.1%; Score 33; DB 4; Length 15;
  Query Match
  Best Local Similarity
                          53.8%; Pred. No. 2e+02;
                                                                             0;
  Matches
            7; Conservative
                                1; Mismatches
                                                  5; Indels
                                                                 0; Gaps
Qу
            2 QTPQPLLQVMMEP 14
              1 1111 :1 1
Db
            3 QPPQPLPPTVMFP 15
RESULT 16
AAB72563
ΙD
     AAB72563 standard; peptide; 15 AA.
XX
```

```
AC
     AAB72563;
XX
     09-MAY-2001 (first entry)
DT
XX
DE
     Colostrinin peptide #32.
XX
     Neuroprotective; neural cell differentiation regulator; colostrinin;
KW
KW
     colostrum.
XX
     Unidentified.
os
XX
PN
     WO200112651-A2.
XX
PD
     22-FEB-2001.
XX
PF
     17-AUG-2000; 2000WO-US022774.
XX
PR
     17-AUG-1999;
                    99US-0149633P.
XX
     (TEXA ) UNIV TEXAS SYSTEM.
PA
XX
ΡI
     Boldogh I;
XX
     WPI; 2001-226545/23.
DR
XX
     Use of colostrinin, its constituent peptide or analog as a neural cell
PT
     regulator, for promoting neural cell differentiation and treating damaged
PT
     neural cells in a patient.
PT
XX
     Claim 6; Page 22; 35pp; English.
PS
XX
     The present invention relates to a method for promoting neural cell
CC
     differentiation and treating damaged neural cells, using colostrinin and
CC
     colostrinin constituent peptides (e.g. the present peptide) as a neural
CC
CC
     cell regulator. Colostrinin is a polypeptide complex found in colostrum
XX
     Sequence 15 AA;
SQ
  Query Match
                          37.1%; Score 33; DB 4; Length 15;
                          53.8%; Pred. No. 2e+02;
  Best Local Similarity
  Matches
             7; Conservative
                                 1; Mismatches
                                                 5; Indels
                                                                 0; Gaps
                                                                              0;
            2 QTPQPLLQVMMEP 14
Qу
              1 1111
                     : 1 1
            3 QPPQPLPPTVMFP 15
Db
RESULT 17
AA014610
     AAO14610 standard; peptide; 15 AA.
ID
XX
     AA014610;
AC
XX
DT
     27-MAY-2002 (first entry)
XX
DE
     Neural cell regulatory colostrinin peptide 32.
XX
```

```
KW
     Neural cell differentiation; neural cell regulator; colostrinin peptide;
KW
     neural cell formation; proline-rich polypeptide aggregate; colostrum;
KW
     neural cell treatment.
XX
    Unidentified.
os
XX
FH
                     Location/Qualifiers
    Key
FT
    Modified-site
FT
                     /note= "Optional C-terminal amide"
XX
    WO200213851-A1.
PN
XX
PD
     21-FEB-2002.
XX
    17-AUG-2000; 2000WO-US022777.
PF
XX
PR
    17-AUG-2000; 2000WO-US022777.
XX
PΑ
     (TEXA ) UNIV TEXAS SYSTEM.
XX
PΙ
    Boldogh I, Stanton JG, Hughes TK;
XX
DR
    WPI; 2002-269152/31.
XX
PT
     Promoting cell differentiation in a patient involves use of blood cell
PT
     regulator selected from colostrinin, its constituent peptide and/or
PT
     analog.
XX
PS
    Claim 7; Page 22; 37pp; English.
XX
    The invention comprises a method for promoting cell differentiation (e.g.
CC
CC
    neural cell differentiation). The method involves contacting cells with a
CC
    neural cell regulator (i.e. a colostrinin peptide) in order to change the
CC
     cells in morphology to form neural cells. Colostrinin is a proline-rich
     polypeptide aggregate that is present in colostrum. The method of the
CC
CC
     invention is useful for promoting the differentiation of cells and for
CC
    treating damaged neural cells in a patient. The present amino acid
CC
     sequence represents a specifically claimed colostrinin peptide used in
CC
     the method of the invention
XX
SQ
     Sequence 15 AA;
  Query Match
                          37.1%;
                                  Score 33; DB 5; Length 15;
                          53.8%; Pred. No. 2e+02;
  Best Local Similarity
            7; Conservative
 Matches
                                 1; Mismatches
                                                                              0;
                                                   5; Indels
                                                                 0; Gaps
            2 QTPQPLLQVMMEP 14
Qу
              1 1111 :11
Db
            3 QPPQPLPPTVMFP 15
RESULT 18
AAM51066
     AAM51066 standard; peptide; 15 AA.
XX
AC
    AAM51066;
XX
```

```
DT
     30-MAY-2002 (first entry)
XX
DE
     Colostrinin constituent peptide (casein amino acids 159-173).
XX
KW
     Colostrinin; colostrum; immunomodulator; cardiovascular;
     blood cell regulator; cytokine inducer; beta-casein; human.
KW
XX
os
     Homo sapiens.
XX
FH
                     Location/Qualifiers
     Key
FT
    Modified-site
FT
                     /note= "optional C-terminal amidation"
XX
PN
    WO200213849-A1.
XX
PD
     21-FEB-2002.
XX
PF
     17-AUG-2000; 2000WO-US022775.
XX
     17-AUG-2000; 2000WO-US022775.
PR
XX
PA
     (TEXA ) UNIV TEXAS SYSTEM.
     (REGE-) REGEN THERAPEUTICS PLC.
PΑ
XX
     Stanton GJ, Hughes TK, Boldogh I, Georgiades J;
ΡI
XX
     WPI; 2002-269150/31.
DR
XX
     Modulation of blood cell proliferation in a patient involves use of blood
PT
PT
     cell regulator selected from colostrinin, its constituent peptide and/or
PΤ
     analog.
XX
     Claim 1; Page 34; 54pp; English.
PS
XX
CC
     The present sequence is that of a colostrinin constituent peptide that is
CC
     used as an immunological regulator and as a blood cell regulator in
     claimed methods of the invention. It is classified as having a beta-
CC
CC
     casein homologue precursor, and corresponds to casein amino acids 159-
CC
     173. Methods are claimed for: inducing a cytokine in a cell by contact
CC
     with an immunological regulator, where the cell is present in a cell
CC
     culture, a tissue, an organ or an organism, and the cell is mammalian,
CC
     including human; modulating an immune response in a cell by contact with
     the immunological regulator under conditions effective to induce a
CC
     cytokine; modulating an immune response in a patient by administering an
CC
     immunological regulator under conditions effective to induce a cytokine,
CC
CC
     where the immunological regulator is administered topically or as part of
CC
     a dietary supplement, and where the immune response is specific or non
CC
     specific, an interferon response or an antibody response; modulating
CC
     blood cell proliferation by contacting blood cells with a blood cell
CC
     regulator, where the blood cells are present in a cell culture or an
CC
     organism, are mammalian or human, and where the blood cells are increased
     in number or differentiated; and a method for modulating blood cell
CC
CC
     proliferation in a patent. A claimed cytokine-inducing composition
     comprises a pharmaceutical carrier and an active agent such as the
CC
CC
     present peptide
XX
```

SQ

Sequence 15 AA;

```
37.1%;
  Query Match
                                  Score 33; DB 5; Length 15;
  Best Local Similarity
                          53.8%; Pred. No. 2e+02;
             7; Conservative
                                 1: Mismatches
                                                   5; Indels
                                                                  0; Gaps
                                                                              0;
Qу
            2 QTPQPLLQVMMEP 14
              1 1111
                      : 1 1
Db
            3 QPPQPLPPTVMFP 15
RESULT 19
AAE20261
     AAE20261 standard; peptide; 15 AA.
XX
AC
    AAE20261;
XX
DT
    18-JUN-2002
                 (first entry)
XX
DΕ
    Colostrinin constituent peptide #32.
XX
KW
     Blood cell regulator; colostrinin; constituent peptide; oxidative stress;
KW
     therapy; oxidative damage; skin; aging; wound healing; cell replacement;
KW
     tissue; organ; cosmetic procedure; repair; regeneration; preservation;
KW
     transplantation; implantation; dermatological; vulnerary.
XX
os
     Unidentified.
XX
FΗ
     Key
                     Location/Qualifiers
FT
    Modified-site
FT
                     /note= "Optionally C-terminal amide"
XX
PN
     WO200213850-A1.
XX
PD
     21-FEB-2002.
XX
     17-AUG-2000; 2000WO-US022776.
PF
XX
PR
     17-AUG-2000; 2000WO-US022776.
XX
     (TEXA ) UNIV TEXAS SYSTEM.
PA
XX
PΙ
     Stanton GJ, Hughes TK, Boldogh I;
XX
DR
     WPI; 2002-269151/31.
XX
PT
     Composition useful for the modulation of blood cell proliferation in a
PT
     patient comprises a blood cell regulator selected from colostrinin, its
PT
     constituent peptide and/or analog.
XX
PS
     Claim 6; Page 26; 51pp; English.
XX
CC
     The invention relates to a composition which comprises a blood cell
CC
     regulator selected from colostrinin, its constituent peptide and/or
CC
     analogue. The invention is used for modulating the oxidative stress level
CC
     in a cell e.g. mammalian or human cell present in a cell culture, tissue,
CC
     organ, or organism; or for treating oxidative damage to the skin of a
CC
     patient e.g. animal or human; to modulate oxidative stress during/ after
```

```
CC
    a premature birth or normal birth, preventing/delaying aging in a
CC
    patient, enhancing wound healing, and the reduction of side effects of
CC
    cosmetic procedures. The method changes the level of an oxidising species
CC
    in the cell, such as decreases or prevents increase in the level of
CC
    damage to a biomolecule of the patient selected from DNA, protein and/or
CC
    lipid, compared to the same conditions when the oxidative stress
CC
    regulator is not present. The modulation of oxidative stress results in
CC
    enhanced repair, regeneration, and replacement of cells, tissues and
CC
    organs (e.g. kidney, liver, pancreas, skin, and the other internal and
    external organs), as well as enhanced preservation of such organs for
CC
CC
    transplantation, implantation, or scientific research. The present
CC
    sequence is a colostrinin constituent peptide
XX
SQ
    Sequence 15 AA;
 Query Match
                          37.1%; Score 33; DB 5; Length 15;
 Best Local Similarity
                         53.8%; Pred. No. 2e+02;
 Matches
                              1; Mismatches
            7; Conservative
                                                  5; Indels
                                                                0; Gaps
                                                                            0;
           2 QTPQPLLQVMMEP 14
Qу
             Db
           3 QPPQPLPPTVMFP 15
RESULT 20
AAR21055
ID
    AAR21055 standard; peptide; 8 AA.
XX
AC
    AAR21055;
XX
DT
    25-MAR-2003
                 (revised)
    01-JUN-1992 (first entry)
DT
XX
DE
    Gamma-carboxylase, N-terminus.
XX
KW
    Degenerate; Vitamin K dependent proteins; PCR.
XX
os
    Homo sapiens.
XX
PN
    WO9201795-A.
XX
PD
    06-FEB-1992.
XX
PF
    23-JUL-1990;
                   90US-00557220.
XX
PR
    23-JUL-1990;
                   90US-00557220.
PR
    14-MAR-1991;
                   91US-00669735.
XX
    (ZYMO ) ZYMOGENETICS INC.
PA
XX
PΙ
    Berkner KL;
XX
DR
    WPI; 1992-064951/08.
XX
PT
    Gamma-carboxylase protein compsns. - used in recombinant prodn. of active
PT
    vitamin=K dependent proteins.
XX
```

```
PS
     Claim 6; Table 9; 91pp; English.
XX
     Nucleotide sequences encoding gamma-carboxylase were obtd. using PCR, and
CC
     oligonucleotides designed from amino acid sequences determined by
CC
CC
    microsequencing of partially purified material. SEQ ID No 19 (peptide 5)
CC
     is one of seven alternative sequences for the N-terminus of gamma
CC
     carboxylase, due to the degeneracy of the genetic code. Obtaining the
CC
     full DNA and protein sequence of gamma-carboxylase will allow proteins
     such as Factor VII, Factor IX, prothrombin, (activated) protein C,
CC
CC
     protein S, protein Z, or osteocalcin to be easily produced by recombinant
CC
     techniques. See also AAR21049-56, AAR23010. (Updated on 25-MAR-2003 to
CC
     correct PA field.)
XX
SO
     Sequence 8 AA;
  Query Match
                          36.0%; Score 32; DB 2; Length 8;
  Best Local Similarity
                          100.0%; Pred. No. 1.4e+06;
                                0; Mismatches
                                                  0; Indels
  Matches
            6; Conservative
                                                                 0; Gaps
                                                                              0;
            3 TPQPLL 8
Qy
              111111
Db
            1 TPQPLL 6
RESULT 21
ABR31148
    ABR31148 standard; peptide; 15 AA.
ID
XX
AC
    ABR31148;
XX
\mathsf{DT}
     19-MAY-2003 (first entry)
XX
     Human cancer-related protein 83P4B8 HLA peptide #2210.
DE
XX
KW
     Human; cytostatic; vaccine; cancer; immune response; HLA;
KW
     human leukocyte antigen.
XX
os
     Homo sapiens.
XX
PN
     WO200283921-A2.
XX
PD
     24-OCT-2002.
XX
PF
     10-APR-2002; 2002WO-US011654.
XX
PR
     10-APR-2001; 2001US-0282739P.
     10-APR-2001; 2001US-0283112P.
PR
     25-APR-2001; 2001US-0286630P.
PR
XX
     (AGEN-) AGENSYS INC.
PΑ
XX
PΙ
     Jakobovits A, Challita-Eid PM, Faris M, Ge W, Hubert RS;
ΡI
     Morrison K, Morrison RK, Raitano AB;
XX
DR
     WPI; 2003-075555/07.
XX
PT
     New composition comprising a substance that modulates the structure of
```

```
proteins and polynucleotides, useful for therapeutic, prognostic and
PΤ
PТ
     diagnostic reagents for eliciting cellular or humoral immune response in
PΤ
     cancer patients.
XX
PS
    Claim 13; Page 487; 1021pp; English.
XX
CC
    The present invention relates to novel human cancer-related genes and
CC
    proteins (ABZ78120-ABZ78168 and ABR01789-ABR01861). The genes and
    proteins are useful for eliciting a humoral or cellular immune response.
CC
CC
    The genes are useful as probes and primers for the amplification and/or
CC
    detection of genes, mRNAs or their fragments, as reagents for the
CC
     diagnosis and/or prognosis of cancer, as coding sequences capable of
     directing the expression of the protein, as tools for modulating or
CC
     inhibiting the expression of genes and/or translation of transcripts, and
CC
CC
     as therapeutic agents. The proteins and peptides are useful as
CC
     therapeutic, prognostic and diagnostic reagents for cancer. The present
CC
     sequence is a human leukocyte antigen (HLA) peptide, used in an example
    from the invention
CC
XX
SQ
     Sequence 15 AA;
                          36.0%; Score 32; DB 6; Length 15;
  Query Match
  Best Local Similarity
                          58.3%; Pred. No. 2.9e+02;
             7; Conservative
                                2; Mismatches
 Matches
                                                  3; Indels
                                                                 0; Gaps
                                                                              0;
            1 LQTPQPLLQVMM 12
Qу
              1: ||||:||
            4 LKAVQPLLKVSM 15
RESULT 22
ABR31149
    ABR31149 standard; peptide; 15 AA.
ID
XX
AC
    ABR31149;
XX
DT
    19-MAY-2003
                 (first entry)
XX
DE
     Human cancer-related protein 83P4B8 HLA peptide #2211.
XX
KW
     Human; cytostatic; vaccine; cancer; immune response; HLA;
KW
     human leukocyte antigen.
XX
OS
     Homo sapiens.
XX
PN
     WO200283921-A2.
XX
     24-OCT-2002.
PD
XX
     10-APR-2002; 2002WO-US011654.
PF
XX
PR
     10-APR-2001; 2001US-0282739P.
PR
     10-APR-2001; 2001US-0283112P.
PR
     25-APR-2001; 2001US-0286630P.
XX
PA
     (AGEN-) AGENSYS INC.
XX
```

```
PΙ
     Jakobovits A, Challita-Eid PM, Faris M, Ge W, Hubert RS;
ΡI
     Morrison K, Morrison RK, Raitano AB;
XX
     WPI; 2003-075555/07.
DR
XX
PT
     New composition comprising a substance that modulates the structure of
PT
     proteins and polynucleotides, useful for therapeutic, prognostic and
PT
     diagnostic reagents for eliciting cellular or humoral immune response in
PT
     cancer patients.
XX
PS
     Claim 13; Page 487; 1021pp; English.
XX
CC
     The present invention relates to novel human cancer-related genes and
CC
     proteins (ABZ78120-ABZ78168 and ABR01789-ABR01861). The genes and
     proteins are useful for eliciting a humoral or cellular immune response.
CC
CC
     The genes are useful as probes and primers for the amplification and/or
CC
     detection of genes, mRNAs or their fragments, as reagents for the
CC
     diagnosis and/or prognosis of cancer, as coding sequences capable of
CC
     directing the expression of the protein, as tools for modulating or
CC
     inhibiting the expression of genes and/or translation of transcripts, and
CC
     as therapeutic agents. The proteins and peptides are useful as
CC
     therapeutic, prognostic and diagnostic reagents for cancer. The present
CC
     sequence is a human leukocyte antigen (HLA) peptide, used in an example
CC
     from the invention
XX
SQ
     Sequence 15 AA;
  Query Match
                          36.0%; Score 32; DB 6; Length 15;
                          58.3%; Pred. No. 2.9e+02;
  Best Local Similarity
  Matches
            7; Conservative
                                 2; Mismatches
                                                3; Indels
                                                                             0;
                                                                 0; Gaps
            1 LOTPOPLLOVMM 12
Qу
             1: ||||:||
Db
            1 LKAVQPLLKVSM 12
RESULT 23
ABR30440
     ABR30440 standard; peptide; 15 AA.
XX
AC
     ABR30440;
XX
DΤ
     19-MAY-2003 (first entry)
XX
DE
     Human cancer-related protein 83P4B8 HLA peptide #1502.
XX
KW
     Human; cytostatic; vaccine; cancer; immune response; HLA;
KW
     human leukocyte antigen.
XX
os
     Homo sapiens.
XX
PN
     WO200283921-A2.
XX
PD
     24-OCT-2002.
XX
PF
     10-APR-2002; 2002WO-US011654.
XX
```

```
PR
     10-APR-2001; 2001US-0282739P.
PR
     10-APR-2001; 2001US-0283112P.
PR
     25-APR-2001; 2001US-0286630P.
XX
PA
     (AGEN-) AGENSYS INC.
XX
PΙ
     Jakobovits A, Challita-Eid PM, Faris M, Ge W, Hubert RS;
PΙ
     Morrison K, Morrison RK, Raitano AB;
XX
DR
     WPI; 2003-075555/07.
XX
PT
     New composition comprising a substance that modulates the structure of
PT
     proteins and polynucleotides, useful for therapeutic, prognostic and
PT
     diagnostic reagents for eliciting cellular or humoral immune response in
PT
     cancer patients.
XX
PS
     Claim 13; Page 475; 1021pp; English.
XX
CC
     The present invention relates to novel human cancer-related genes and
CC
     proteins (ABZ78120-ABZ78168 and ABR01789-ABR01861). The genes and
     proteins are useful for eliciting a humoral or cellular immune response.
CC
CC
     The genes are useful as probes and primers for the amplification and/or
CC
     detection of genes, mRNAs or their fragments, as reagents for the
CC
     diagnosis and/or prognosis of cancer, as coding sequences capable of
     directing the expression of the protein, as tools for modulating or
CC
CC
     inhibiting the expression of genes and/or translation of transcripts, and
CC
     as therapeutic agents. The proteins and peptides are useful as
CC
     therapeutic, prognostic and diagnostic reagents for cancer. The present
     sequence is a human leukocyte antigen (HLA) peptide, used in an example
CC
CC
     from the invention
XX
SQ
     Sequence 15 AA;
  Query Match
                          36.0%;
                                 Score 32; DB 6; Length 15;
  Best Local Similarity
                          58.3%; Pred. No. 2.9e+02;
  Matches
            7; Conservative
                              2; Mismatches 3; Indels
                                                                 0; Gaps
                                                                             0;
            1 LQTPQPLLQVMM 12
Qу
             1: ||||:|||
Db
            1 LKAVQPLLKVSM 12
RESULT 24
ABR31517
     ABR31517 standard; peptide; 15 AA.
ID
XX
AC
    ABR31517;
XX
DT
     19-MAY-2003 (first entry)
XX
DE
     Human cancer-related protein 83P4B8 HLA peptide #2579.
XX
KW
     Human; cytostatic; vaccine; cancer; immune response; HLA;
KW
     human leukocyte antigen.
XX
OS
     Homo sapiens.
XX
```

```
PN
     WO200283921-A2.
XX
PD
     24-OCT-2002.
XX
PF
     10-APR-2002; 2002WO-US011654.
XX
PR
     10-APR-2001; 2001US-0282739P.
     10-APR-2001; 2001US-0283112P.
PR
PR
     25-APR-2001; 2001US-0286630P.
XX
PA
     (AGEN-) AGENSYS INC.
XX
PΙ
     Jakobovits A, Challita-Eid PM, Faris M, Ge W, Hubert RS;
ΡI
     Morrison K, Morrison RK, Raitano AB;
XX
DR
     WPI; 2003-075555/07.
XX
     New composition comprising a substance that modulates the structure of
PT
     proteins and polynucleotides, useful for therapeutic, prognostic and
PT
PT
     diagnostic reagents for eliciting cellular or humoral immune response in
PT
     cancer patients.
XX
PS
     Claim 13; Page 494; 1021pp; English.
XX
CC
     The present invention relates to novel human cancer-related genes and
CC
     proteins (ABZ78120-ABZ78168 and ABR01789-ABR01861). The genes and
CC
     proteins are useful for eliciting a humoral or cellular immune response.
CC
     The genes are useful as probes and primers for the amplification and/or
     detection of genes, mRNAs or their fragments, as reagents for the
CC
     diagnosis and/or prognosis of cancer, as coding sequences capable of
CC
CC
     directing the expression of the protein, as tools for modulating or
CC
     inhibiting the expression of genes and/or translation of transcripts, and
CC
     as therapeutic agents. The proteins and peptides are useful as
CC
     therapeutic, prognostic and diagnostic reagents for cancer. The present
CC
     sequence is a human leukocyte antigen (HLA) peptide, used in an example
CC
     from the invention
XX
SQ
     Sequence 15 AA;
  Query Match
                          36.0%; Score 32; DB 6; Length 15;
  Best Local Similarity 58.3%; Pred. No. 2.9e+02;
                               2; Mismatches 3; Indels
  Matches
                                                                 0; Gaps
                                                                             0;
             7; Conservative
            1 LQTPQPLLQVMM 12
Qу
              1: 1111:11
Db
            1 LKAVQPLLKVSM 12
RESULT 25
ABR30536
     ABR30536 standard; peptide; 15 AA.
XX
AC
     ABR30536;
XX
     19-MAY-2003 (first entry)
DT
XX
DE
     Human cancer-related protein 83P4B8 HLA peptide #1598.
```

```
XX
KW
    Human; cytostatic; vaccine; cancer; immune response; HLA;
KW
    human leukocyte antigen.
XX
OS
    Homo sapiens.
XX
    WO200283921-A2.
PN
XX
     24-OCT-2002.
PD
XX
     10-APR-2002; 2002WO-US011654.
PF
XX
    10-APR-2001; 2001US-0282739P.
PR
PR
    10-APR-2001; 2001US-0283112P.
PR
     25-APR-2001; 2001US-0286630P.
XX
PA
     (AGEN-) AGENSYS INC.
XX
PΙ
    Jakobovits A, Challita-Eid PM, Faris M, Ge W, Hubert RS;
ΡI
    Morrison K, Morrison RK, Raitano AB;
XX
DR
    WPI; 2003-075555/07.
XX
PT
    New composition comprising a substance that modulates the structure of
    proteins and polynucleotides, useful for therapeutic, prognostic and
PT
PT
    diagnostic reagents for eliciting cellular or humoral immune response in
PT
     cancer patients.
XX
PS
    Claim 13; Page 476; 1021pp; English.
XX
CC
    The present invention relates to novel human cancer-related genes and
CC
    proteins (ABZ78120-ABZ78168 and ABR01789-ABR01861). The genes and
CC
    proteins are useful for eliciting a humoral or cellular immune response.
    The genes are useful as probes and primers for the amplification and/or
CC
CC
    detection of genes, mRNAs or their fragments, as reagents for the
CC
    diagnosis and/or prognosis of cancer, as coding sequences capable of
CC
    directing the expression of the protein, as tools for modulating or
CC
     inhibiting the expression of genes and/or translation of transcripts, and
CC
     as therapeutic agents. The proteins and peptides are useful as
CC
     therapeutic, prognostic and diagnostic reagents for cancer. The present
CC
     sequence is a human leukocyte antigen (HLA) peptide, used in an example
CC
     from the invention
XX
SQ
     Sequence 15 AA;
  Query Match
                          36.0%; Score 32; DB 6; Length 15;
  Best Local Similarity
                          58.3%; Pred. No. 2.9e+02;
  Matches
            7; Conservative
                                2; Mismatches
                                                  3; Indels
                                                                 0; Gaps
                                                                             0;
            1 LOTPOPLLOVMM 12
Qy
              1: ||||:||
            4 LKAVOPLLKVSM 15
Db
RESULT 26
```

RESULT 26
ABR30847
ID ABR30847 standard; peptide; 15 AA.

```
XX
AC
    ABR30847;
XX
DT
     19-MAY-2003 (first entry)
XX
DE
     Human cancer-related protein 83P4B8 HLA peptide #1909.
XX
KW
     Human; cytostatic; vaccine; cancer; immune response; HLA;
KW
     human leukocyte antigen.
XX
OS
     Homo sapiens.
XX
    WO200283921-A2.
PN
XX
PD
    24-OCT-2002.
XX
PF
    10-APR-2002; 2002WO-US011654.
XX
PR
     10-APR-2001; 2001US-0282739P.
     10-APR-2001; 2001US-0283112P.
PR
     25-APR-2001; 2001US-0286630P.
PR
XX
PA
     (AGEN-) AGENSYS INC.
XX
     Jakobovits A, Challita-Eid PM, Faris M, Ge W, Hubert RS;
PΙ
    Morrison K, Morrison RK, Raitano AB;
PI
XX
    WPI; 2003-075555/07.
DR
XX
     New composition comprising a substance that modulates the structure of
PT
     proteins and polynucleotides, useful for therapeutic, prognostic and
PΤ
PT
     diagnostic reagents for eliciting cellular or humoral immune response in
PΤ
     cancer patients.
XX
PS
     Claim 13; Page 482; 1021pp; English.
XX
CC
     The present invention relates to novel human cancer-related genes and
CC
     proteins (ABZ78120-ABZ78168 and ABR01789-ABR01861). The genes and
CC
     proteins are useful for eliciting a humoral or cellular immune response.
CC
     The genes are useful as probes and primers for the amplification and/or
CC
     detection of genes, mRNAs or their fragments, as reagents for the
CC
     diagnosis and/or prognosis of cancer, as coding sequences capable of
     directing the expression of the protein, as tools for modulating or
CC
CC
     inhibiting the expression of genes and/or translation of transcripts, and
CC
     as therapeutic agents. The proteins and peptides are useful as
CC
     therapeutic, prognostic and diagnostic reagents for cancer. The present
CC
     sequence is a human leukocyte antigen (HLA) peptide, used in an example
CC
     from the invention
XX
SO
     Sequence 15 AA;
                          36.0%; Score 32; DB 6; Length 15;
  Query Match
  Best Local Similarity
                          58.3%; Pred. No. 2.9e+02;
            7: Conservative
                                2; Mismatches
                                                 3; Indels
                                                                 0; Gaps
                                                                              0;
            1 LQTPQPLLQVMM 12
Qу
```

1: ||||:|| |

CC

CC

CC

```
RESULT 27
AAB23189
     AAB23189 standard; peptide; 12 AA.
XX
AC
     AAB23189;
XX
DT
     29-JAN-2001 (first entry)
XX
     Hsp47-binding hydrophobic phage display dodecapeptide, SEQ ID NO:11.
DE
XX
     Hsp47-binding hydrophobic dodecapeptide; phage display library;
KW
KW
     Hsp47 external domain; carcinoma; cancer; targetting molecule; therapy;
KW
     diagnosis; detection; imaging; drug delivery; invasion; migration;
KW
    metastasis; modulation; tumour; skin; basal cells; colon;
KW
     large intestine; lung; breast; bladder; oral cancer;
KW
     head and neck cancer; larynx; nasopharynx; adrenal cortex;
KW
     apocrine gland; kidney; liver; pancreas; prostate.
XX
os
     Synthetic.
XX
PN
    WO200054805-A1.
XX
PD
     21-SEP-2000.
XX
PF
     15-MAR-2000; 2000WO-US006588.
XX
PR
     15-MAR-1999;
                    99US-0124481P.
XX
     (UYMA-) UNIV MARYLAND BALTIMORE.
PA
XX
PΙ
     Sauk JJ;
XX
DR
    WPI; 2000-655997/63.
XX
PT
    Treating, diagnosing or modulating a carcinoma cell, which expresses Heat
PT
     shock protein 47 on its surface, involves administering an agent
PT
     comprising targeting moiety which binds to Hsp47 external domain.
XX
PS
     Claim 6; Page 8; 87pp; English.
XX
CC
     The invention relates to methods of treating and diagnosing carcinomas in
     which heat shock protein 47 (Hsp47) is expressed on the surface of the
CC
CC
     carcinoma cells, involving administering an agent comprising a targetting
CC
    moiety which specifically binds to the external domain of Hsp47. The
CC
     invention also relates to peptides (AAB23181-B23203) which specifically
    bind to external domains of such surface-localised Hsp47 molecules and
CC
CC
     have sequences encompassed by the generic sequences XHyHyXXHyXXXXHyHy or
CC
     HyXXXHyHyXXHyXXX where X, independently, can be any amino acid and Hy,
CC
     independently, can be any hydrophobic amino acid. The invention also
CC
     encompasses methods of screening for agents which bind Hsp47 external
CC
     domains. Hsp47-binding agents can be may be used to treat Hsp47-
```

expressing carcinomas, and for modulating the activity of a tumour cell

interaction with the extracellular matrix. The targetting moiety (such as

with respect to invasion, migration, motility or metastasis, or to its

```
CC
     an Hsp47-binding peptide, a toxin or an antibody) may be coupled with a
CC
     therapeutic moiety (such as a cytotoxic agent or a therapeutic gene) for
CC
     cancer treatment, or with a detectable moiety for imaging. Carcinomas
CC
     which may be treated or diagnosed according to methods of the invention
CC
     include those of the skin, basal cells, large intestine, lung, colon,
CC
    breast, bladder, oral, head and neck, larynx, nasopharynx, adrenal
CC
     cortex, apocrine glands, kidney, liver, pancreas, or prostate. Targetting
CC
     carcinoma cells with Hsp47-binding agents results in efficient delivery
     of therapeutic agents, reduced doses, reduced side effects and sensitive
CC
CC
     detection or imaging of carcinoma cells. Sequences AAB23181-B23191
CC
     represent predominantly hydrophobic phage display library dodecapeptides
     identified as being able to bind Hsp47 in an exmplification of the
CC
CC
     invention
XX
     Sequence 12 AA;
SQ
  Query Match
                          34.8%;
                                  Score 31; DB 3; Length 12;
  Best Local Similarity
                         41.7%; Pred. No. 3.3e+02;
                                 3; Mismatches
  Matches
            5; Conservative
                                                   4; Indels
                                                                 0; Gaps
                                                                             0;
           3 TPQPLLQVMMEP 14
Qу
             Db
           1 TPQPNMMLRISP 12
RESULT 28
AAM97279
     AAM97279 standard; peptide; 14 AA.
ID
XX
AC
    AAM97279;
XX
DT
     24-JAN-2002 (first entry)
XX
DE
    Human peptide #554 encoded by a SNP oligonucleotide.
XX
     Immunosuppressive; immunostimulatory; antiinflammatory; cytostatic;
KW
KW
     neuroprotective; antimicrobial; gene therapy; vaccine; amylase; cancer;
KW
     amyloid protein; angiopoietin; apoptosis related protein; cadherin;
KW
     cyclin; polymerase; oncogene; histone; kinase; colony stimulating factor;
     complement related protein; cytochrome; kinesin; cytokine; interferon;
KW
KW
     interleukin; G-protein coupled receptor; thioesterase; inflammation;
KW
    multifactorial disease; autoimmune disease; infection;
KW
     nervous system disease.
XX
OS
    Homo sapiens.
XX
PN
    WO200147944-A2.
XX
     05-JUL-2001.
PD
XX
     28-DEC-2000; 2000WO-US035498.
PF
XX
PR
     28-DEC-1999;
                    99US-0173419P.
PR
     27-DEC-2000; 2000US-00173419.
XX
PA
     (CURA-) CURAGEN CORP.
XX
```

```
ΡI
     Shimkets RA, Leach M;
XX
DR
     WPI; 2001-465210/50.
XX
PT
     Polymorphic nucleic acids encoding e.g. amylases, cyclins, polymerases,
PT
     oncogenes and histones, useful for diagnosing and treating, e.g. cancer,
PT
     autoimmune diseases and infections.
XX
PS
     Disclosure; Page 3789; 4143pp; English.
XX
CC
     The present invention relates to oligonucleotides (see AAL26793-AAL34659)
CC
     encoding polymorphic variants of proteins related to amylases, amyloid
CC
     proteins, angiopoietin, apoptosis related proteins, cadherin, cyclin,
CC
     polymerase, oncogenes, histones, kinases, colony stimulating factors,
CC
     complement related proteins, cytochromes, kinesins, cytokines,
CC
     interferons, interleukins, G-protein coupled receptors and thioesterases.
CC
     The present sequence is a peptide encoded by one such oligonucleotide.
CC
     The oligonucleotides and the peptides encoded by them may be used in the
CC
     prevention, diagnosis and treatment of diseases associated with
CC
     inappropriate expression of the proteins listed above. Disorders that may
CC
     be prevented, diagnosed and/or treated include multifactorial diseases
CC
     with a genetic component, such as autoimmune diseases (e.g. rheumatoid
CC
     arthritis, multiple sclerosis, diabetes, systemic lupus erythromatosus
CC
     and Grave's disease), inflammation, cancer (e.g. cancers of the bladder,
CC
     brain, breast, colon and kidney, leukaemia), diseases of the nervous
CC
     system and an infection of pathogenic organisms
XX
SO
     Sequence 14 AA;
                          34.8%;
  Query Match
                                  Score 31; DB 4; Length 14;
  Best Local Similarity
                          75.0%; Pred. No. 3.9e+02;
                                                  1; Indels
  Matches
             6; Conservative
                                 1; Mismatches
                                                                  0; Gaps
                                                                              0;
            6 PLLQVMME 13
Qу
              11:1 111
Db
            1 PLIQEMME 8
RESULT 29
AAW56352
ID
     AAW56352 standard; peptide; 15 AA.
XX
AC
     AAW56352;
XX
DT
     17-AUG-1998 (first entry)
XX
DE
     LO-CD2a light chain framework region 2.
XX
KW
     LO-CD2a; monoclonal antibody; CD2; rat; chimeric antibody;
KW
     humanised antibody; antibody engineering; graft rejection;
     graft versus host disease; autoimmune disease; therapy.
KW
XX
OS
     Rattus sp.
XX
FΗ
                     Location/Qualifiers
     Kev
FT
     Misc-difference 2
FT
                     /note= "retained in humanised antibody"
```

```
FT
    Misc-difference 3
FT
                     /note= "retained in humanised antibody"
FT
    Misc-difference 11
FT
                     /note= "retained in humanised antibody"
FT
    Misc-difference 12
FT
                     /note= "retained in humanised antibody"
XX
PN
    WO9807444-A1.
XX
PD
     26-FEB-1998.
XX
PF
     16-AUG-1996;
                    96WO-US013281.
XX
PR
     16-AUG-1996;
                    96WO-US013281.
XX
PΑ
     (BIOT-) BIOTRANSPLANT INC.
PΑ
     (UYLO-) UNIV CATHOLIQUE LOUVAIN.
XX
ΡI
     Bazin H, Latinne D, Kaplan R, Kieber-Emmons T, Postema CE;
ΡI
    White-Scharf ME;
XX
DR
    WPI; 1998-168898/15.
XX
PT
     Humanised antibody - comprises complementarity determining region from LO
PT
     -CD2a, useful to prevent or inhibit graft versus host or auto-immune
PT
     disease.
XX
PS
     Example 7; Page 58; 133pp; English.
XX
CC
     This peptide comprises framework region 2 (FR 2) of the light chain (see
CC
     AAW56347) of rat anti-CD2 monoclonal antibody LO-CD2a. FRs 1-4 (see
     AAW56351-54) were compared with FRs 1-4 of the human HUM5400 light chain.
CC
CC
     A novel humanized LO-CD2a light chain (see AAW56349) comprises rat LO-
     CD2a complementarity determining regions and HUM5400 constant regions,
CC
     but with amino acids 9, 12, 41, 42, 50, 51 and 82 of the rat framework
CC
     retained to maintain binding specificity. The chimeric antibody can be
CC
     used to inhibit an immune response, specifically human T cell activation
CC
CC
     and proliferation, in a patient, and to inhibit rejection of a graft in a
CC
     patient (claimed), useful for preventing or inhibiting graft versus host
CC
     or autoimmune disease
XX
SQ
     Sequence 15 AA;
                          34.8%; Score 31; DB 2; Length 15;
  Query Match
                          71.4%; Pred. No. 4.1e+02;
  Best Local Similarity
  Matches
             5; Conservative
                              2; Mismatches
                                                  0; Indels
                                                                 0; Gaps
                                                                              0;
            2 QTPQPLL 8
Qу
              1:111:
Db
            8 OSPOPLI 14
RESULT 30
AAY30191
     AAY30191 standard; peptide; 15 AA.
ID
XX
AC
    AAY30191;
```

```
XX
DT
     29-OCT-1999 (first entry)
XX
     Framework 2 region of rat monoclonal antibody LO-CD2a light chain.
DΕ
XX
     Antibody LO-CD2a; CD2 antigen; T-lymphocyte; humanised antibody;
KW
KW
     T-cell-mediated immune response; graft rejection; autoimmune disease;
     graft-versus-host disease; T cell; natural killer cell.
KW
XX
os
     Rattus sp.
XX
PN
    US5951983-A.
XX
PD
     14-SEP-1999.
XX
ΡF
     07-JUN-1995;
                    95US-00477989.
XX
                    93US-00027008.
PR
     05-MAR-1993;
     09-SEP-1993;
                    93US-00119032.
PR
PR
     29-MAR-1995;
                    95US-00407009.
XX
     (BIOT-) BIO TRANSPLANT INC.
PΑ
PA
     (UYLO-) UNIV CATHOLIQUE LOUVAIN.
XX
     White-Scharf ME, Postema CE, Kaplan R, Latinne D, Bazin H;
PΙ
PΙ
     Kieber-Emmons T;
XX
DR
     WPI; 1999-526991/44.
XX
PT
     Antibody mediated Inhibition of T cell immune response.
XX
PS
     Example 7; Col 33-34; 104pp; English.
XX
CC
     The present sequence represent a framework region of rat monoclonal
CC
     antibody LO-CD2a. LO-CD2a binds to an epitope of a CD2 antigen T-
CC
     lymphocytes. The present sequence is compared to the corresponding region
CC
     from a human V kappa gene, and used to design humanised sequences. The
CC
     humanized LO-CD2a antibody comprises the human constant regions, a light
     chain framework region derived from a human antibody, a heavy chain
CC
CC
     framework region derived from a human antibody, heavy and light chain
CC
     complementarity determining regions (CDRs) of the non-human monoclonal
     antibody produced by the cell line deposited as ATCC HB11423. The
СC
CC
     humanised antibodies are used in a method for treating a patient to
     inhibit a T-cell-mediated immune response. The method is useful for the
CC
     treatment or prevention of graft rejection and graft-versus-host disease,
CC
CC
     as well as in the treatment of autoimmune diseases which are mediated by
CC
     the activation and proliferation of T cells or natural killer cells
XX
SO
     Sequence 15 AA;
                          34.8%; Score 31; DB 2; Length 15;
  Query Match
                          71.4%; Pred. No. 4.1e+02;
  Best Local Similarity
  Matches
             5; Conservative 2; Mismatches 0; Indels
                                                                     Gaps
Qу
            2 QTPQPLL 8
              1:1111:
Db
            8 QSPQPLI 14
```

```
RESULT 31
AAU74511
ID
     AAU74511 standard; peptide; 15 AA.
XX
AC
     AAU74511;
XX
DT
     09-APR-2002
                 (first entry)
XX
DE
     Human ATP dependent serine proteinase 21 N-terminal peptide.
XX
     Human; ATP dependent serine protease 21; malignant tumour; HIV;
KW
KW
     haemopathy; human immunodeficiency virus; immunological disease;
KW
     inflammation; cytostatic; haemostatic; virucide; immunomodulatory;
KW
     antiinflammatory; gene therapy.
XX
OS
     Homo sapiens.
XX
PN
     WO200194529-A2.
XX
PD
     13-DEC-2001.
XX
     28-APR-2001; 2001WO-CN000658.
PF
XX
PR
     29-APR-2000; 2000CN-00115512.
XX
PA
     (SHAN-) SHANGHAI BIOWINDOW GENE DEV INC.
XX
ΡI
     Mao Y, Xie Y;
XX
     WPI; 2002-090143/12.
DR
XX
PT
     Human ATP dependent serien proteinase 21 and encoding polynucleotide,
     used in diagnosis and treatment of malignant tumors, hemopathy, human
PT
     immunodeficiency virus infection, immunological diseases and
PT
РΤ
     inflammation.
XX
PS
     Example 5; Page 18; 32pp; Chinese.
XX
CC
     The invention relates to an isolated polypeptide of human ATP dependent
CC
     serine proteinase 21. The protein of the invention and its associated
CC
     cDNA sequence are used in the treatment of various diseases including
CC
     malignant tumours, haemopathy, human immunodeficiency virus (HIV)
CC
     infection, immunological diseases and various inflammations. This
CC
     sequence represents the N-terminal peptide of ATP dependent serine
CC
     proteinase 21, used in ELISA
XX
SO
     Sequence 15 AA;
  Query Match
                          34.8%;
                                  Score 31; DB 5; Length 15;
  Best Local Similarity
                          46.2%;
                                  Pred. No. 4.1e+02;
  Matches
                                 3; Mismatches 4; Indels
             6; Conservative
                                                                  0;
                                                                               0;
                                                                      Gaps
            5 QPLLQVMMEPQGD 17
Qу
              | | :::|||
Db
            3 QLLESTVIQPQGD 15
```

```
RESULT 32
ABR38577
ΙD
     ABR38577 standard; peptide; 15 AA.
XX
AC
     ABR38577;
XX
DT
     19-MAY-2003
                 (first entry)
XX
DE
     Human cancer-related protein 192P2G7 HLA peptide #1437.
XX
KW
     Human; cytostatic; vaccine; cancer; immune response; HLA;
KW
     human leukocyte antigen.
XX
OS
     Homo sapiens.
XX
PN
     W0200283921-A2.
XX
PD
     24-OCT-2002.
XX
     10-APR-2002; 2002WO-US011654.
PF
XX
PR
     10-APR-2001; 2001US-0282739P.
     10-APR-2001; 2001US-0283112P.
PR
     25-APR-2001; 2001US-0286630P.
₽R
XX
PΑ
     (AGEN-) AGENSYS INC.
XX
PΙ
     Jakobovits A, Challita-Eid PM, Faris M, Ge W, Hubert RS;
PΙ
     Morrison K, Morrison RK, Raitano AB;
XX
DR
     WPI; 2003-075555/07.
XX
PT
     New composition comprising a substance that modulates the structure of
     proteins and polynucleotides, useful for therapeutic, prognostic and
PT
PT
     diagnostic reagents for eliciting cellular or humoral immune response in
PT
     cancer patients.
XX
PS
     Claim 13; Page 622; 1021pp; English.
XX
CC
     The present invention relates to novel human cancer-related genes and
     proteins (ABZ78120-ABZ78168 and ABR01789-ABR01861). The genes and
CC
     proteins are useful for eliciting a humoral or cellular immune response.
CC
     The genes are useful as probes and primers for the amplification and/or
CC
CC
     detection of genes, mRNAs or their fragments, as reagents for the
CC
     diagnosis and/or prognosis of cancer, as coding sequences capable of
CC
     directing the expression of the protein, as tools for modulating or
CC
     inhibiting the expression of genes and/or translation of transcripts, and
     as therapeutic agents. The proteins and peptides are useful as
CC
CC
     therapeutic, prognostic and diagnostic reagents for cancer. The present
CC
     sequence is a human leukocyte antigen (HLA) peptide, used in an example
CC
     from the invention
XX
SO
     Sequence 15 AA;
```

```
Best Local Similarity
                          46.2%; Pred. No. 4.1e+02;
  Matches 6; Conservative
                                3; Mismatches
                                                4; Indels
                                                                 0; Gaps
                                                                             0;
           1 LQTPQPLLQVMME 13
Qу
              1: | | | | :: |
            2 LEYPQPGLDIIKE 14
Db
RESULT 33
ABR38568
     ABR38568 standard; peptide; 15 AA.
ΙD
XX
AC
    ABR38568;
XX
DT
     19-MAY-2003 (first entry)
XX
DE
     Human cancer-related protein 192P2G7 HLA peptide #1428.
XX
KW
     Human; cytostatic; vaccine; cancer; immune response; HLA;
KW
     human leukocyte antigen.
XX
OS
     Homo sapiens.
XX
PN
     WO200283921-A2.
XX
PD
     24-OCT-2002.
XX
     10-APR-2002; 2002WO-US011654.
PF
XX
     10-APR-2001; 2001US-0282739P.
PR
     10-APR-2001; 2001US-0283112P.
PR
     25-APR-2001; 2001US-0286630P.
PR
XX
PA
     (AGEN-) AGENSYS INC.
XX
PΙ
     Jakobovits A, Challita-Eid PM, Faris M, Ge W, Hubert RS;
PΙ
     Morrison K, Morrison RK, Raitano AB;
XX
     WPI; 2003-075555/07.
DR
XX
PT
     New composition comprising a substance that modulates the structure of
PT
     proteins and polynucleotides, useful for therapeutic, prognostic and
PT
     diagnostic reagents for eliciting cellular or humoral immune response in
PT
     cancer patients.
XX
PS
     Claim 13; Page 622; 1021pp; English.
XX
CC
     The present invention relates to novel human cancer-related genes and
     proteins (ABZ78120-ABZ78168 and ABR01789-ABR01861). The genes and
CC
CC
     proteins are useful for eliciting a humoral or cellular immune response.
CC
     The genes are useful as probes and primers for the amplification and/or
     detection of genes, mRNAs or their fragments, as reagents for the
CC
CC
     diagnosis and/or prognosis of cancer, as coding sequences capable of
     directing the expression of the protein, as tools for modulating or
CC
CC
     inhibiting the expression of genes and/or translation of transcripts, and
CC
     as therapeutic agents. The proteins and peptides are useful as
CC
     therapeutic, prognostic and diagnostic reagents for cancer. The present
```

```
CC
     sequence is a human leukocyte antigen (HLA) peptide, used in an example
CC
     from the invention
XX
SO
     Sequence 15 AA;
  Query Match
                          34.8%; Score 31; DB 6; Length 15;
  Best Local Similarity
                         46.2%; Pred. No. 4.1e+02;
  Matches
            6; Conservative 3; Mismatches 4; Indels
                                                                 0; Gaps
                                                                             0;
           1 LQTPQPLLQVMME 13
Qy
             1: | | | | :: |
Db
           3 LEYPQPGLDIIKE 15
RESULT 34
ABR38729
     ABR38729 standard; peptide; 15 AA.
XX
AC
    ABR38729;
XX
DT
    19-MAY-2003 (first entry)
XX
DE
     Human cancer-related protein 192P2G7 HLA peptide #1589.
XX
ΚW
     Human; cytostatic; vaccine; cancer; immune response; HLA;
KW
     human leukocyte antigen.
XX
os
     Homo sapiens.
XX
PN
     WO200283921-A2.
XX
PD
    24-OCT-2002.
XX
     10-APR-2002; 2002WO-US011654.
PF
XX
     10-APR-2001; 2001US-0282739P.
PR
PR
     10-APR-2001; 2001US-0283112P.
PR
     25-APR-2001; 2001US-0286630P.
XX
PA
     (AGEN-) AGENSYS INC.
XX
     Jakobovits A, Challita-Eid PM, Faris M, Ge W, Hubert RS;
ΡI
PΙ
     Morrison K, Morrison RK, Raitano AB;
XX
DR
     WPI; 2003-075555/07.
XX
PT
     New composition comprising a substance that modulates the structure of
     proteins and polynucleotides, useful for therapeutic, prognostic and
PT
PT
     diagnostic reagents for eliciting cellular or humoral immune response in
PT
     cancer patients.
XX
PS
     Claim 13; Page 624; 1021pp; English.
XX
CC
     The present invention relates to novel human cancer-related genes and
     proteins (ABZ78120-ABZ78168 and ABR01789-ABR01861). The genes and
CC
CC
     proteins are useful for eliciting a humoral or cellular immune response.
CC
     The genes are useful as probes and primers for the amplification and/or
```

```
CC
     detection of genes, mRNAs or their fragments, as reagents for the
CC
     diagnosis and/or prognosis of cancer, as coding sequences capable of
CC
     directing the expression of the protein, as tools for modulating or
CC
     inhibiting the expression of genes and/or translation of transcripts, and
CC
     as therapeutic agents. The proteins and peptides are useful as
CC
     therapeutic, prognostic and diagnostic reagents for cancer. The present
СC
     sequence is a human leukocyte antigen (HLA) peptide, used in an example
CC
     from the invention
XX
SO
     Sequence 15 AA;
  Query Match
                          34.8%; Score 31; DB 6; Length 15;
  Best Local Similarity
                         46.2%; Pred. No. 4.1e+02;
  Matches
            6; Conservative
                                3; Mismatches 4; Indels
                                                                 0; Gaps
                                                                             0;
           1 LQTPQPLLQVMME 13
Qу
             1: ||| | :: |
Db
           2 LEYPQPGLDIIKE 14
RESULT 35
ADD15523
    ADD15523 standard; peptide; 10 AA.
ΙD
XX
AC
    ADD15523;
XX
DT
     15-JAN-2004 (first entry)
XX
DΕ
    Predicted epitopic region of the S aureus GlmM protein (SeqID 88).
XX
KW
    microbial; antimicrobial; membrane biosynthesis; pathogenic;
KW
     immunological response; vaccination; surface disinfectant;
KW
    personal hygiene application; food preservative; GlmM;
KW
    phosphoglucosamine-mutase; epitope.
XX
     Staphylococcus aureus.
os
XX
PN
    WO2003025007-A2.
XX
PD
     27-MAR-2003.
XX
     20-SEP-2002; 2002WO-CA001428.
PF
XX
     21-SEP-2001; 2001US-0323992P.
PR
     21-SEP-2001; 2001US-0324152P.
PR
     25-SEP-2001; 2001US-0324692P.
PR
     26-OCT-2001; 2001US-0339924P.
     29-OCT-2001; 2001US-0350973P.
PR
     30-OCT-2001; 2001US-0340924P.
PR
     27-NOV-2001; 2001US-0333666P.
PR
    18-DEC-2001; 2001US-0341732P.
PR
PR
     18-DEC-2001; 2001US-0341776P.
PR
     19-DEC-2001; 2001US-0341949P.
XX
PA
     (AFFI-) AFFINIUM PHARM INC.
XX
ΡI
     Edwards A, Dharamsi A, Vedadi M, Alam MZ, Awrey D, Beattie B;
```

```
PΙ
     Domagala M,
                 Houston S, Kanagarajah D, Nethery K, Ng I, Mansoury K;
PΙ
    Mcdonald M,
                 Pinder B, Viola C, Wrezel O;
XX
DR
    WPI; 2003-468119/44.
XX
PT
    Novel crystallized recombinant polypeptides from Staphylococcus aureus,
PT
     Streptococcus pneumoniae and Escherichia coli and which are involved in
PT
    membrane biosynthesis, useful as targets for pathogenic bacteria.
XX
PS
     Disclosure; SEQ ID NO 88; 325pp; English.
XX
    This invention relates to the structural and functional characterisation
CC
CC
     of microbial polypeptides from Staphylococcus aureus (S. aureus),
CC
     Streptococcus pneumoniae (S. pneumoniae) and Escherichia coli (E. coli)
CC
     that provide novel antimicrobial targets. Specifically, it refers to
CC
    polypeptides that are involved in membrane biosynthesis, which play a
CC
    critical role in the life cycle and viability of their pathogenic species
CC
    of origin, and hence provide valuable drug targets. Furthermore, the
CC
     invention describes modified version of these proteins that facilitate
CC
     characterisation by labelling with isotopic or heavy atoms, and also
CC
     fusion proteins. These proteins provide structural and functional
CC
     information to aid the discovery of therapeutic molecules to treat
CC
     disorders associated with a particular pathogenic species. As such, they
CC
     are useful for inducing an immunological response in an individual and as
CC
     an antigen for vaccination purposes. The polypeptides are also useful for
CC
     developing antimicrobial agents for use as surface disinfectants,
CC
     personal hygiene applications and as food preservatives or in treating
CC
     food products to eliminate potential pathogens. This peptide sequence is
CC
     a predicted epitopic sequence of the S. aureus GlmM protein of the
CC
    invention.
XX
     Sequence 10 AA;
SO
  Query Match
                          33.7%;
                                  Score 30; DB 7; Length 10;
                          55.6%; Pred. No. 3.9e+02;
  Best Local Similarity
                                                                 0; Gaps
             5; Conservative
                                 4; Mismatches
                                                0; Indels
                                                                             0;
            5 QPLLQVMME 13
Qу
              : | | : : | | : |
Db
            2 EPLVRVMVE 10
RESULT 36
AAM96749
ID
    AAM96749 standard; peptide; 14 AA.
XX
AC
    AAM96749;
XX
DT
    24-JAN-2002 (first entry)
XX
DE
    Human peptide #24 encoded by a SNP oligonucleotide.
XX
KW
     Immunosuppressive; immunostimulatory; antiinflammatory; cytostatic;
KW
     neuroprotective; antimicrobial; gene therapy; vaccine; amylase; cancer;
KW
     amyloid protein; angiopoietin; apoptosis related protein; cadherin;
     cyclin; polymerase; oncogene; histone; kinase; colony stimulating factor;
KW
KW
     complement related protein; cytochrome; kinesin; cytokine; interferon;
```

```
KW
     interleukin; G-protein coupled receptor; thioesterase; inflammation;
KW
     multifactorial disease; autoimmune disease; infection;
KW
     nervous system disease.
XX
os
     Homo sapiens.
XX
PN
     WO200147944-A2.
XX
PD
     05-JUL-2001.
XX
     28-DEC-2000; 2000WO-US035498.
PF
XX
     28-DEC-1999;
                    99US-0173419P.
PR
     27-DEC-2000; 2000US-00173419.
PR
XX
PA
     (CURA-) CURAGEN CORP.
XX
PΙ
     Shimkets RA, Leach M;
XX
DR
     WPI; 2001-465210/50.
XX
PT
     Polymorphic nucleic acids encoding e.g. amylases, cyclins, polymerases,
PΤ
     oncogenes and histones, useful for diagnosing and treating, e.g. cancer,
     autoimmune diseases and infections.
PT
XX
     Disclosure; Page 3673; 4143pp; English.
PS
XX
CC
     The present invention relates to oligonucleotides (see AAL26793-AAL34659)
     encoding polymorphic variants of proteins related to amylases, amyloid
CC
     proteins, angiopoietin, apoptosis related proteins, cadherin, cyclin,
CC
CC
     polymerase, oncogenes, histones, kinases, colony stimulating factors,
     complement related proteins, cytochromes, kinesins, cytokines,
CC
CC
     interferons, interleukins, G-protein coupled receptors and thioesterases.
     The present sequence is a peptide encoded by one such oligonucleotide.
CC
CC
     The oligonucleotides and the peptides encoded by them may be used in the
     prevention, diagnosis and treatment of diseases associated with
CC
     inappropriate expression of the proteins listed above. Disorders that may
CC
CC
     be prevented, diagnosed and/or treated include multifactorial diseases
CC
     with a genetic component, such as autoimmune diseases (e.g. rheumatoid
CC
     arthritis, multiple sclerosis, diabetes, systemic lupus erythromatosus
CC
     and Grave's disease), inflammation, cancer (e.g. cancers of the bladder,
     brain, breast, colon and kidney, leukaemia), diseases of the nervous
CC
CC
     system and an infection of pathogenic organisms
XX
SQ
     Sequence 14 AA;
                          33.7%;
                                  Score 30; DB 4; Length 14;
  Query Match
  Best Local Similarity
                          46.2%; Pred. No. 5.6e+02;
  Matches
             6; Conservative 4; Mismatches
                                                  3; Indels
                                                                 0; Gaps
                                                                              0;
Qy
            4 POPLLOVMMEPOG 16
              1:|| :|:: ||
            1 PRPLRRVVLFYQG 13
Db
```

```
AAM47320 standard; peptide; 15 AA.
ID
XX
     AAM47320;
AC
XX
DT
     22-FEB-2002 (first entry)
XX
DΕ
     Human zinc finger structural domain 52 N-terminal peptide.
XX
KW
     Human; zinc finger structural domain 52; cancer; nosohaemia;
KW
     immunological disease; HIV infection; inflammation; gene therapy.
XX
os
     Homo sapiens.
XX
PN
     CN1306987-A.
XX
PD
     08-AUG-2001.
XX
PF
     28-JAN-2000; 2000CN-00111572.
XX
     28-JAN-2000; 2000CN-00111572.
PR
XX
     (BODA-) BODAO GENE TECH CO LTD SHANGHAI.
PA
XX
PΙ
     Mao Y, Xie Y;
XX
DR
     WPI; 2002-026871/04.
XX
PΤ
     Polypeptide-zinc finger structural domain 52 and polynucleotide for
PΤ
     coding said polypeptide.
XX
PS
     Example 6; Page 18(Disclosure); 35pp; Chinese.
XX
CC
     The present invention provides the protein and coding sequences of human
     zinc finger structural domain 52. The sequences can be used in the
CC
     treatment of cancer, nosohaemia, HIV infection, immunological diseases
CC
CC
     and inflammation. The present sequence is the N-terminus of the protein
CC
     of the invention
XX
SO
     Sequence 15 AA;
  Query Match
                          33.7%; Score 30; DB 5; Length 15;
                          42.9%; Pred. No. 6e+02;
  Best Local Similarity
             6; Conservative
                                 2; Mismatches
  Matches
                                                   6; Indels
                                                                  0; Gaps
                                                                              0;
            1 LQTPQPLLQVMMEP 14
Qу
              ::|| || |
Db
            1 METPTPLPPVPASP 14
RESULT 38
ABR30169
ID
     ABR30169 standard; peptide; 15 AA.
XX
AC
     ABR30169;
XX
DT
     19-MAY-2003 (first entry)
XX
```

```
DE
     Human cancer-related protein 74P3B3 HLA peptide #3013.
XX
KW
     Human; cytostatic; vaccine; cancer; immune response; HLA;
KW
     human leukocyte antigen.
XX
os
     Homo sapiens.
XX
PN
     W0200283921-A2.
XX
PD
     24-OCT-2002.
XX
     10-APR-2002; 2002WO-US011654.
PF
XX
PR
     10-APR-2001; 2001US-0282739P.
PR
     10-APR-2001; 2001US-0283112P.
     25-APR-2001; 2001US-0286630P.
PR
XX
     (AGEN-) AGENSYS INC.
PA
XX
ΡI
     Jakobovits A, Challita-Eid PM, Faris M, Ge W, Hubert RS;
PΙ
    Morrison K, Morrison RK, Raitano AB;
XX
DR
     WPI; 2003-075555/07.
XX
PT
     New composition comprising a substance that modulates the structure of
     proteins and polynucleotides, useful for therapeutic, prognostic and
PT
PT
     diagnostic reagents for eliciting cellular or humoral immune response in
PT
     cancer patients.
XX
     Claim 13; Page 470; 1021pp; English.
PS
XX
CC
     The present invention relates to novel human cancer-related genes and
     proteins (ABZ78120-ABZ78168 and ABR01789-ABR01861). The genes and
CC
     proteins are useful for eliciting a humoral or cellular immune response.
CC
CC
     The genes are useful as probes and primers for the amplification and/or
CC
     detection of genes, mRNAs or their fragments, as reagents for the
CC
     diagnosis and/or prognosis of cancer, as coding sequences capable of
CC
     directing the expression of the protein, as tools for modulating or
CC
     inhibiting the expression of genes and/or translation of transcripts, and
CC
     as therapeutic agents. The proteins and peptides are useful as
CC
     therapeutic, prognostic and diagnostic reagents for cancer. The present
     sequence is a human leukocyte antigen (HLA) peptide, used in an example
CC
CC
     from the invention
XX
     Sequence 15 AA;
SQ
                          33.7%; Score 30; DB 6; Length 15;
  Query Match
                          45.5%; Pred. No. 6e+02;
  Best Local Similarity
                                 2; Mismatches
                                                                              0;
  Matches
             5; Conservative
                                                   4; Indels
                                                                 0; Gaps
            4 PQPLLQVMMEP 14
Qу
             | |: | :||
Db
            5 PPPIKQCSLEP 15
```

```
ΙD
     ABR30262 standard; peptide; 15 AA.
XX
AC
    ABR30262;
XX
DT
     19-MAY-2003 (first entry)
XX
DE
     Human cancer-related protein 74P3B3 HLA peptide #3106.
XX
KW
     Human; cytostatic; vaccine; cancer; immune response; HLA;
KW
     human leukocyte antigen.
XX
os
     Homo sapiens.
XX
PN
     WO200283921-A2.
XX
PD
     24-OCT-2002.
XX
PF
     10-APR-2002; 2002WO-US011654.
XX
PR
     10-APR-2001; 2001US-0282739P.
     10-APR-2001; 2001US-0283112P.
PR
PR
     25-APR-2001; 2001US-0286630P.
XX
PΑ
     (AGEN-) AGENSYS INC.
XX
PΙ
     Jakobovits A, Challita-Eid PM, Faris M, Ge W, Hubert RS;
PΙ
     Morrison K, Morrison RK, Raitano AB;
XX
     WPI; 2003-075555/07.
DR
XX
     New composition comprising a substance that modulates the structure of
PT
PT
     proteins and polynucleotides, useful for therapeutic, prognostic and
PT
     diagnostic reagents for eliciting cellular or humoral immune response in
PT
     cancer patients.
XX
PS
     Claim 13; Page 471; 1021pp; English.
XX
CC
     The present invention relates to novel human cancer-related genes and
CC
     proteins (ABZ78120-ABZ78168 and ABR01789-ABR01861). The genes and
CC
     proteins are useful for eliciting a humoral or cellular immune response.
CC
     The genes are useful as probes and primers for the amplification and/or
CC
     detection of genes, mRNAs or their fragments, as reagents for the
CC
     diagnosis and/or prognosis of cancer, as coding sequences capable of
CC
     directing the expression of the protein, as tools for modulating or
CC
     inhibiting the expression of genes and/or translation of transcripts, and
CC
     as therapeutic agents. The proteins and peptides are useful as
CC
     therapeutic, prognostic and diagnostic reagents for cancer. The present
     sequence is a human leukocyte antigen (HLA) peptide, used in an example
CC
CC
     from the invention
XX
SQ
     Sequence 15 AA;
                          33.7%; Score 30; DB 6; Length 15;
  Query Match
  Best Local Similarity
                          45.5%; Pred. No. 6e+02;
                                                                 0; Gaps
  Matches
             5; Conservative
                                 2; Mismatches 4; Indels
                                                                              0;
```

```
RESULT 40
ABR30274
ID
     ABR30274 standard; peptide; 15 AA.
XX
AC
     ABR30274;
XX
DΤ
     19-MAY-2003 (first entry)
XX
DE
     Human cancer-related protein 74P3B3 HLA peptide #3118.
XX
KW
     Human; cytostatic; vaccine; cancer; immune response; HLA;
KW
     human leukocyte antigen.
XX
OS
    Homo sapiens.
XX
PN
    WO200283921-A2.
XX
PD
     24-OCT-2002.
XX
PF
     10-APR-2002; 2002WO-US011654.
XX
     10-APR-2001; 2001US-0282739P.
PR
     10-APR-2001; 2001US-0283112P.
PR
     25-APR-2001; 2001US-0286630P.
PR
XX
PA
     (AGEN-) AGENSYS INC.
XX
PΙ
     Jakobovits A, Challita-Eid PM, Faris M, Ge W, Hubert RS;
PΙ
    Morrison K, Morrison RK, Raitano AB;
XX
DR
     WPI; 2003-075555/07.
XX
PT
     New composition comprising a substance that modulates the structure of
PТ
     proteins and polynucleotides, useful for therapeutic, prognostic and
PT
     diagnostic reagents for eliciting cellular or humoral immune response in
PT
     cancer patients.
XX
PS
     Claim 13; Page 472; 1021pp; English.
XX
CC
     The present invention relates to novel human cancer-related genes and
CC
     proteins (ABZ78120-ABZ78168 and ABR01789-ABR01861). The genes and
CC
     proteins are useful for eliciting a humoral or cellular immune response.
CC
     The genes are useful as probes and primers for the amplification and/or
CC
     detection of genes, mRNAs or their fragments, as reagents for the
CC
     diagnosis and/or prognosis of cancer, as coding sequences capable of
     directing the expression of the protein, as tools for modulating or
CC
CC
     inhibiting the expression of genes and/or translation of transcripts, and
CC
     as therapeutic agents. The proteins and peptides are useful as
CC
     therapeutic, prognostic and diagnostic reagents for cancer. The present
CC
     sequence is a human leukocyte antigen (HLA) peptide, used in an example
CC
     from the invention
XX
SO
     Sequence 15 AA;
```

```
Query Match
                          33.7%; Score 30; DB 6; Length 15;
  Best Local Similarity
                         45.5%; Pred. No. 6e+02;
                                                                 0; Gaps
  Matches
             5; Conservative
                                 2; Mismatches
                                                   4; Indels
                                                                              0;
            4 PQPLLQVMMEP 14
Qу
             1 1: 1 :11
            5 PPPIKQCSLEP 15
Db
RESULT 41
ABR30229
ID
     ABR30229 standard; peptide; 15 AA.
XX
AC
    ABR30229;
XX
DT
     19-MAY-2003 (first entry)
XX
DE
     Human cancer-related protein 74P3B3 HLA peptide #3073.
XX
KW
     Human; cytostatic; vaccine; cancer; immune response; HLA;
KW
     human leukocyte antigen.
XX
OS
     Homo sapiens.
XX
     WO200283921-A2.
PN
XX
     24-OCT-2002.
PD
XX
     10-APR-2002; 2002WO-US011654.
PF
XX
     10-APR-2001; 2001US-0282739P.
PR
     10-APR-2001; 2001US-0283112P.
PR
     25-APR-2001; 2001US-0286630P.
PR
XX
PΑ
     (AGEN-) AGENSYS INC.
XX
     Jakobovits A, Challita-Eid PM,
PΙ
                                      Faris M,
                                                Ge W, Hubert RS;
     Morrison K, Morrison RK, Raitano AB;
PΙ
XX
     WPI; 2003-075555/07.
DR
XX
     New composition comprising a substance that modulates the structure of
PТ
     proteins and polynucleotides, useful for therapeutic, prognostic and
PT
     diagnostic reagents for eliciting cellular or humoral immune response in
PΤ
PT
     cancer patients.
XX
     Claim 13; Page 471; 1021pp; English.
PS
XX
CC
     The present invention relates to novel human cancer-related genes and
     proteins (ABZ78120-ABZ78168 and ABR01789-ABR01861). The genes and
CC
     proteins are useful for eliciting a humoral or cellular immune response.
CC
     The genes are useful as probes and primers for the amplification and/or
CC
     detection of genes, mRNAs or their fragments, as reagents for the
CC
CC
     diagnosis and/or prognosis of cancer, as coding sequences capable of
CC
     directing the expression of the protein, as tools for modulating or
CC
     inhibiting the expression of genes and/or translation of transcripts, and
```

```
CC
     as therapeutic agents. The proteins and peptides are useful as
CC
     therapeutic, prognostic and diagnostic reagents for cancer. The present
CC
     sequence is a human leukocyte antigen (HLA) peptide, used in an example
CC
     from the invention
XX
SQ
     Sequence 15 AA;
  Query Match
                          33.7%; Score 30; DB 6; Length 15;
  Best Local Similarity
                          45.5%; Pred. No. 6e+02;
             5; Conservative
                                2; Mismatches 4; Indels
                                                                 0; Gaps
                                                                              0;
            4 PQPLLQVMMEP 14
Qy
              1 1: 1 : 1 1
Db
            4 PPPIKQCSLEP 14
RESULT 42
ABR30154
     ABR30154 standard; peptide; 15 AA.
XX
AC
    ABR30154;
XX
DT
    19-MAY-2003 (first entry)
XX
DE
     Human cancer-related protein 74P3B3 HLA peptide #2998.
XX
KW
     Human; cytostatic; vaccine; cancer; immune response; HLA;
KW
     human leukocyte antigen.
XX
OS
     Homo sapiens.
XX
PN
     WO200283921-A2.
XX
PD
    24-OCT-2002.
XX
PF
     10-APR-2002; 2002WO-US011654.
XX
PR
     10-APR-2001; 2001US-0282739P.
PR
     10-APR-2001; 2001US-0283112P.
PR
     25-APR-2001; 2001US-0286630P.
XX
PA
     (AGEN-) AGENSYS INC.
XX
PΙ
     Jakobovits A, Challita-Eid PM, Faris M, Ge W, Hubert RS;
PΙ
     Morrison K, Morrison RK, Raitano AB;
XX
DR
     WPI; 2003-075555/07.
XX
PT
     New composition comprising a substance that modulates the structure of
PT
     proteins and polynucleotides, useful for therapeutic, prognostic and
PT
     diagnostic reagents for eliciting cellular or humoral immune response in
PT
     cancer patients.
XX
PS
     Claim 13; Page 469; 1021pp; English.
XX
CC
     The present invention relates to novel human cancer-related genes and
CC
     proteins (ABZ78120-ABZ78168 and ABR01789-ABR01861). The genes and
```

```
CC
     proteins are useful for eliciting a humoral or cellular immune response.
CC
    The genes are useful as probes and primers for the amplification and/or
CC
     detection of genes, mRNAs or their fragments, as reagents for the
CC
     diagnosis and/or prognosis of cancer, as coding sequences capable of
     directing the expression of the protein, as tools for modulating or
CC
CC
     inhibiting the expression of genes and/or translation of transcripts, and
     as therapeutic agents. The proteins and peptides are useful as
CC
     therapeutic, prognostic and diagnostic reagents for cancer. The present
CC
CC
     sequence is a human leukocyte antigen (HLA) peptide, used in an example
CC
     from the invention
XX
SQ
     Sequence 15 AA;
  Query Match
                          33.7%; Score 30; DB 6; Length 15;
  Best Local Similarity
                          45.5%; Pred. No. 6e+02;
                                2; Mismatches
                                                   4; Indels
 Matches
             5; Conservative
                                                                 0; Gaps
                                                                             0;
            4 PQPLLQVMMEP 14
Qу
             1 1: 1 : 11
Db
            2 PPPIKQCSLEP 12
RESULT 43
ABR31758
    ABR31758 standard; peptide; 15 AA.
ID
XX
AC
    ABR31758;
XX
DT
     19-MAY-2003 (first entry)
XX
DE
    Human cancer-related protein 109P1D4 HLA peptide #1530.
XX
KW
     Human; cytostatic; vaccine; cancer; immune response; HLA;
KW
     human leukocyte antigen.
XX
OS
    Homo sapiens.
XX
PN
     WO200283921-A2.
XX
PD
     24-OCT-2002.
XX
     10-APR-2002; 2002WO-US011654.
PF
XX
     10-APR-2001; 2001US-0282739P.
PR
     10-APR-2001; 2001US-0283112P.
PR
PR
     25-APR-2001; 2001US-0286630P.
XX
     (AGEN-) AGENSYS INC.
PA
XX
     Jakobovits A, Challita-Eid PM, Faris M, Ge W, Hubert RS;
PΙ
PΙ
     Morrison K, Morrison RK, Raitano AB;
XX
     WPI; 2003-075555/07.
DR
XX
PT
     New composition comprising a substance that modulates the structure of
PT
     proteins and polynucleotides, useful for therapeutic, prognostic and
PT
     diagnostic reagents for eliciting cellular or humoral immune response in
```

```
PT
     cancer patients.
XX
PS
    Claim 13; Page 498; 1021pp; English.
XX
CC
     The present invention relates to novel human cancer-related genes and
CC
     proteins (ABZ78120-ABZ78168 and ABR01789-ABR01861). The genes and
CC
     proteins are useful for eliciting a humoral or cellular immune response.
CC
     The genes are useful as probes and primers for the amplification and/or
CC
     detection of genes, mRNAs or their fragments, as reagents for the
CC
    diagnosis and/or prognosis of cancer, as coding sequences capable of
     directing the expression of the protein, as tools for modulating or
CC
     inhibiting the expression of genes and/or translation of transcripts, and
CC
CC
     as therapeutic agents. The proteins and peptides are useful as
CC
     therapeutic, prognostic and diagnostic reagents for cancer. The present
     sequence is a human leukocyte antigen (HLA) peptide, used in an example
CC
CC
     from the invention
XX
SQ
    Sequence 15 AA;
                          33.7%; Score 30; DB 6; Length 15;
  Query Match
  Best Local Similarity
                          45.5%; Pred. No. 6e+02;
 Matches
            5; Conservative
                                2; Mismatches
                                                  4; Indels
                                                                 0; Gaps
                                                                             0;
            3 TPQPLLQVMME 13
Qу
              : | | | | : |
Db
            5 SPQPAFQIQPE 15
RESULT 44
ABR30297
    ABR30297 standard; peptide; 15 AA.
XX
AC
    ABR30297;
XX
DT
    19-MAY-2003 (first entry)
XX
DE
    Human cancer-related protein 74P3B3 HLA peptide #3141.
XX
KW
     Human; cytostatic; vaccine; cancer; immune response; HLA;
KW
     human leukocyte antigen.
XX
OS
     Homo sapiens.
XX
    WO200283921-A2.
PN
XX
PD
    24-OCT-2002.
XX
    10-APR-2002; 2002WO-US011654.
PF
XX
     10-APR-2001; 2001US-0282739P.
PR
PR
     10-APR-2001; 2001US-0283112P.
PR
     25-APR-2001; 2001US-0286630P.
XX
PA
     (AGEN-) AGENSYS INC.
XX
PΙ
     Jakobovits A, Challita-Eid PM, Faris M,
                                                Ge W, Hubert RS;
PΙ
     Morrison K, Morrison RK, Raitano AB;
```

```
XX
DR
     WPI; 2003-075555/07.
XX
PT
     New composition comprising a substance that modulates the structure of
PT
     proteins and polynucleotides, useful for therapeutic, prognostic and
     diagnostic reagents for eliciting cellular or humoral immune response in
PT
PT
     cancer patients.
XX
PS
     Claim 13; Page 472; 1021pp; English.
XX
     The present invention relates to novel human cancer-related genes and
CC
     proteins (ABZ78120-ABZ78168 and ABR01789-ABR01861). The genes and
CC
     proteins are useful for eliciting a humoral or cellular immune response.
CC
CC
     The genes are useful as probes and primers for the amplification and/or
CC
     detection of genes, mRNAs or their fragments, as reagents for the
CC
     diagnosis and/or prognosis of cancer, as coding sequences capable of
CC
     directing the expression of the protein, as tools for modulating or
CC
     inhibiting the expression of genes and/or translation of transcripts, and
CC
     as therapeutic agents. The proteins and peptides are useful as
     therapeutic, prognostic and diagnostic reagents for cancer. The present
CC
     sequence is a human leukocyte antigen (HLA) peptide, used in an example
CC
CC
     from the invention
XX
SQ
     Sequence 15 AA;
                          33.7%; Score 30; DB 6; Length 15;
  Query Match
                          45.5%; Pred. No. 6e+02;
  Best Local Similarity
             5; Conservative
                                 2; Mismatches
                                                   4; Indels
                                                                  0; Gaps
                                                                              0;
Qy
            4 PQPLLQVMMEP 14
              | |: | :||
Db
            5 PPPIKOCSLEP 15
RESULT 45
AAY65560
     AAY65560 standard; peptide; 16 AA.
ID
XX
AC
     AAY65560;
XX
DT
     01-FEB-2000 (first entry)
XX
DΕ
     Oestrogen receptor alpha ERE binding peptide 15E.
XX
KW
     Oestrogen receptor; estrogen; estradiol; oestrogen response element; ERE;
KW
     binding; biological activity; fingerprint; molecular braille;
     cellular braille; modulation; tamoxifen; breast cancer; ovarian cancer;
KW
     menopause; osteoporosis; selective oestrogen receptor modulator;
KW
     identification; characterisation; classification.
KW
XX
OS
     Synthetic.
os
     Homo sapiens.
XX
PN
     WO9954728-A2.
XX
PD
     28-OCT-1999.
XX
```

```
PF
     26-MAR-1999;
                    99WO-US006664.
XX
PR
     23-APR-1998;
                    98US-0082756P.
     09-SEP-1998;
                    98US-0099656P.
PR
     08-JAN-1999;
PR
                    99US-0115345P.
XX
     (NOVA-) NOVALON PHARM CORP.
PA
XX
     Paige LA, Hamilton PT, Fowlkes DM, Buehrer B, Barnett T;
PΙ
     Mcdonnell DP, Christensen DJ;
PΙ
XX
DR
     WPI; 2000-013281/01.
XX
PT
     Methods for identifying new receptor modulators, especially estrogen
PT
     modulators to treat tamoxifen refractory breast cancer.
XX
PS
     Example 2.2; Page 161; 219pp; English.
XX
CC
     The present invention describes a method for predicting the biological
     activity of new receptor modulating compounds (I) using novel oligomeric
CC
     peptides (biokeys) which have differential abilities to bind to 2
CC
CC
     different receptor conformations. The method is used to identify new
CC
     drugs that are physiological or pharmacological agonists/antagonists and
CC
     that target various receptors, which are involved in certain disease
CC
     conditions. The system may be used as a primary screening tool to
CC
     identify hits, to classify lead compounds from a drug screen to,
CC
     characterise selective oestrogen receptor modulators (SERMs) in terms of
CC
     agonist and antagonist function and to predict possible clinical effects
CC
     of SERMs such as tissue and receptor specificity. The method can also be
     applied to the fractionation of mixtures of SERMs to determine which
CC
CC
     components are producing agonistic and antagonistic activity. The method
CC
     may be used with other receptors (e.g. progesterone, androgen,
     glucocorticoid, thyroid, vitamin D, beta-adrenergic, dopamine and
CC
     epidermal growth factor, to identify, characterise and classify
CC
CC
     modulators of receptor activity. Peptides comprising a LXXLL motif may be
CC
     used to modulate the oestrogen receptor in treating e.g. breast and
CC
     ovarian cancer and ameliorating the effects of menopause, including
CC
     osteoporosis. AAY65439 to AAY65652 represent oestrogen receptor,
CC
     estradiol receptor and oestrogen response element binding peptides given
CC
     in the exemplification of the present invention. AAZ35740 to AAZ35745
CC
     represent oligonucleotides used in the exemplification of the present
CC
     invention
XX
SQ
     Sequence 16 AA;
                          33.7%; Score 30; DB 3; Length 16;
  Query Match
                         45.5%; Pred. No. 6.4e+02;
  Best Local Similarity
                                2; Mismatches
  Matches
            5; Conservative
                                                  4; Indels 0; Gaps
                                                                             0;
            6 PLLQVMMEPQG 16
Qу
             111 :: 1 1
            5 PLLMALLAPPG 15
Db
RESULT 46
AAU86297
     AAU86297 standard; peptide; 16 AA.
```

XX AC AAU86297; XX DT21-MAY-2002 (first entry) XX DE Oestrogen receptor alpha binding peptide 15E. XX KW Oestrogen receptor; breast cancer; combinatorial peptide library; KW receptor modulating compound. XX OS Synthetic. XX PN WO200204956-A2. XX PD 17-JAN-2002. XX PF11-JUL-2001; 2001WO-US021867. XX PR 12-JUL-2000; 2000US-00614865. PR 21-MAY-2001; 2001US-00860688. XX PΑ (KARO-) KARO BIO USA INC. XX PΙ Fowlkes DM, Barnett TR, Buehrer B; XX DR WPI; 2002-154969/20. XX PTIdentifying receptor-binding peptides comprises screening combinatorial PTpeptide library presented in form of cells each of which coexpress one PΤ peptide member and receptor with signal producing system for reporting РT binding. XX PS Disclosure; Page 144; 175pp; English. XX CC The invention relates to identifying a binding peptide which binds a CC CC comprising screening a combinatorial peptide library presented in the CC CC CC system for reporting binding of the peptide to the receptor. Also CC CC

receptor and which is a member of a combinatorial library of peptides, form of cells which coexpress the receptor or its ligand-binding receptor moiety and one member of the library, together with a signal producing included is a method for predicting the receptor-modulating activity of a compound which modulates the biological activity of a receptor comprising (a) identifying peptides which bind the receptor by the method above, (b) using a number of the peptides to predict the receptor- modulating activity of a compound by (i) providing a panel of identified peptides, where the members differ in their ability to bind to the receptor depending on reference conformations the receptor is in, where the effect of a number of reference substances known to modulate the biological activity of the receptor on the binding of each member of the panel is known and is characterised as a reference fingerprint for each reference substance, (ii) screening a test substance of unknown activity relative to the receptor to determine its effect on the binding of each member of the panel to the receptor, thereby obtaining a test fingerprint for the test substance, (iii) comparing the test fingerprint to the reference fingerprints and (iv) predicting the biological activity of the test substance based on the assumption that its biological activity will be similar to that of reference substances with similar fingerprints. The

CC

```
method is useful for identifying a binding peptide which binds a
CC
     vertebrate, mammalian, preferably human receptor, an intracellular,
CC
CC
     nuclear, oestrogen or androgen receptor. The identified peptides which
     bind to the receptor are useful for predicting the receptor-modulating
CC
CC
     activity of a compound (e.g. ant/agonists). The receptor-binding library
CC
     members are useful in the prediction of the ability of small organic
CC
     molecules, suitable for pharmaceutical use (e.g. in the case of oestrogen
CC
     receptors, for breast cancer treatment), to interact with the receptor.
CC
     The analyte-binding molecules can also be used for in vivo imaging. The
CC
     method has several advantages over whole animal-based assay systems in
CC
     that the same technology can be applied to a variety of different
CC
     receptors, the system can be used for high throughput screening and
CC
     compound characterisation, and gives very distinct patterns for agonists
     and antagonists of receptor activity using very much less protein. The
CC
CC
     present sequence is an oestrogen receptor binding peptide from a
CC
     combinatorial peptide library
XX
SO
     Sequence 16 AA;
  Query Match
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                                  Score 30; DB 5; Length 16;
  Best Local Similarity
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 Matches
            5; Conservative
                                 2; Mismatches
                                                                  0; Gaps
                                                                              0;
                                                   4; Indels
            6 PLLQVMMEPQG 16
Qу
              | \cdot | \cdot | :: | |
Dh
            5 PLLMALLAPPG 15
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XX
AC
    AAU01845;
XX
DT
     07-SEP-2001 (first entry)
XX
DE
     Wheat peptide antagonist for A-gliadin 57-73 QE65 #7.
XX
KW
     Wheat; A-gliadin; 57-75 QE65; coeliac disease; gluten intolerance;
KW
     T-cell binding; antagonist; transglutaminase; transgenic plant.
XX
OS
     Triticum aestivum.
XX
     WO200125793-A2.
PN
XX
PD
     12-APR-2001.
XX
     02-OCT-2000; 2000WO-GB003760.
PF
XX
PR
                    99GB-00023306.
     01-OCT-1999;
XX
PA
     (ISIS-) ISIS INNOVATION LTD.
XX
PΙ
     Anderson RP,
                   Hill AVS,
                              Jewell DP;
XX
     WPI; 2001-300179/31.
DR
XX
```

```
PT
     Diagnosing coeliac disease or susceptibility to the disease in an
     individual, by detecting in vitro or in vivo T cells which bind
PΤ
PT
     immunodominant T cell epitope obtained from naturally occurring homolog
PT
     of gliadin.
XX
PS
     Example 11; Page 58; 107pp; English.
XX
     The sequence represents a gliadin peptide corresponding to A-gliadin 57-
CC
     73 which is naturally polymorphic in that region and is antagonistic to A
CC
     -qliadin 57-73 QE65 interferon gamma ELISpot (not defined) response. The
CC
CC
     peptides of the invention are used to test mammalian (preferably human)
     susceptibility to coeliac disease (gluten intolerance). The peptides are
CC
CC
     contacted with a blood sample and T cell recognition measured, a positive
CC
     T-cell recognition indicating a susceptibility to coeliac disease. The
     peptides are useful for inducing tolerance in an individual and
CC
CC
     antagonists to the peptides are useful for treating or preventing coeliac
CC
     disease in an individual and for producing an antibody specific to them
CC
     or a wild-type sequence. A mutant gliadin protein (or its fragment of 15
CC
     amino acids in length) whose wild-type sequence can be modified by
CC
     transglutaminase to a sequence that comprises the epitope, but which has
CC
     been modified in such a way that it does not contain sequence which can
CC
     be modified by transglutaminase to a sequence that comprise the epitope
CC
     is useful for decreasing the ability of gliadin protein to cause Coeliac
CC
     disease. Nucleic acids encoding proteins antagonistic to the T-cell
CC
     binding of the epitopes are useful for obtaining a transgenic plant cell
CC
     or seed and for the production of a protein. The resultant crop plant is
     useful for obtaining a product of a wheat plant, especially grain, which
CC
CC
     is optionally processed into flour or another grain product. Food
CC
     comprising the antagonistic protein is useful instead of a wild-type
CC
     gliadin
XX
     Sequence 17 AA;
SO
                          33.7%; Score 30; DB 4; Length 17;
  Query Match
                          50.0%; Pred. No. 6.8e+02;
  Best Local Similarity
                                                                 0; Gaps
             6; Conservative
                                 1; Mismatches
                                                   5; Indels
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            4 PQPLLQVMMEPQ 15
Qу
              111 1 : 11
Db
            3 PQPFLPELPYPQ 14
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     AAU01825 standard; peptide; 17 AA.
ID
XX
AC
     AAU01825;
XX
ÐΤ
     07-SEP-2001 (first entry)
XX
DE
     Wheat Gliadin peptide M.
XX
KW
     Wheat; A-gliadin; peptide M; coeliac disease; gluten intolerance;
ΚW
     T-cell binding; antagonist; transglutaminase; transgenic plant.
XX
os
     Triticum aestivum.
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XX

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PN
     WO200125793-A2.
XX
PD
     12-APR-2001.
XX
     02-OCT-2000; 2000WO-GB003760.
PF
XX
PR
     01-OCT-1999;
                    99GB-00023306.
XX
PA
     (ISIS-) ISIS INNOVATION LTD.
XX
ΡI
     Anderson RP, Hill AVS, Jewell DP;
XX
DR
     WPI; 2001-300179/31.
XX
PT
     Diagnosing coeliac disease or susceptibility to the disease in an
     individual, by detecting in vitro or in vivo T cells which bind
PT
PT
     immunodominant T cell epitope obtained from naturally occurring homolog
PT
     of gliadin.
XX
     Example 8; Fig 14; 107pp; English.
PS
XX
CC
     The sequence represents wheat Gliadin peptide M, corresponding to A-
CC
     gliadin 57-73 and containing a natural polymorphism. The peptides of the
     invention are used to test mammalian (preferably human) susceptibility to
CC
CC
     coeliac disease (gluten intolerance). The peptides are contacted with a
     blood sample and T cell recognition measured, a positive T-cell
CC
CC
     recognition indicating a susceptibility to coeliac disease. The peptides
     are useful for inducing tolerance in an individual and antagonists to the
CC
     peptides are useful for treating or preventing coeliac disease in an
CC
CC
     individual and for producing an antibody specific to them or a wild-type
CC
     sequence. A mutant gliadin protein (or its fragment of 15 amino acids in
CC
     length) whose wild-type sequence can be modified by transglutaminase to a
CC
     sequence that comprises the epitope, but which has been modified in such
     a way that it does not contain sequence which can be modified by
CC
CC
     transglutaminase to a sequence that comprise the epitope is useful for
CC
     decreasing the ability of gliadin protein to cause Coeliac disease.
     Nucleic acids encoding proteins antagonistic to the T-cell binding of the
CC
CC
     epitopes are useful for obtaining a transgenic plant cell or seed and for
CC
     the production of a protein. The resultant crop plant is useful for
CC
     obtaining a product of a wheat plant, especially grain, which is
CC
     optionally processed into flour or another grain product. Food comprising
CC
     the antagonistic protein is useful instead of a wild-type gliadin
XX
SQ
     Sequence 17 AA;
  Query Match
                          33.7%; Score 30; DB 4; Length 17;
                          50.0%; Pred. No. 6.8e+02;
  Best Local Similarity
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            6; Conservative
                                1; Mismatches
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  Matches
            4 POPLLOVMMEPO 15
Qу
              Db
            3 PQPFLPQLPYPQ 14
RESULT 49
ABP73085
     ABP73085 standard; peptide; 8 AA.
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XX
AC
     ABP73085;
XX
DT
     17-JUN-2003 (first entry)
XX
DΕ
     Amino acid sequence of an epitope derived from KSHV.
XX
KW
     Epitope; KSHV; CD8 T cell; vaccine; dendritic cell; T lymphocyte.
XX
OS
     Synthetic.
os
     Kaposi's sarcoma herpesvirus.
XX
PN
     WO2003014154-A2.
XX
PD
     20-FEB-2003.
XX
ΡF
     05-AUG-2002; 2002WO-GB003594.
XX
     03-AUG-2001; 2001GB-00019038.
PR
XX
PΑ
     (UNLO ) UNIV COLLEGE LONDON.
XX
PΙ
     Boshoff C;
XX
DR
     WPI; 2003-289909/28.
XX
PT
     New polypeptides and expression vectors with an epitope sequence
PT
     recognized by a CD8 T cell, useful in the manufacture of a vaccine for
PT
     the prophylactic and/or therapeutic treatment of Kaposi's sarcoma
PΤ
     herpesvirus infection.
XX
PS
     Claim 2; Page 49; 58pp; English.
XX
CC
     The present sequence represents an epitope, of a formula given in the
     specification. The epitope is derived from Kaposi's sarcoma herpesvirus
CC
CC
     (KSHV), and is recognised by CD8 T cells. Polypeptides comprising the
CC
     epitope are useful for producing a vaccine against KSHV. They are also
     useful for stimulating dendritic cells or T lymphocytes ex vivo
CC
XX
     Sequence 8 AA;
SQ
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  Query Match
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                          83.3%; Pred. No. 1.4e+06;
  Best Local Similarity
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  Matches
                                 1; Mismatches
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Qу
              111:11
            2 PQPVLQ 7
Db
RESULT 50
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ID
     ABB08361 standard; protein; 9 AA.
XX
AC
     ABB08361;
XX
DT
     07-MAY-2002 (first entry)
```

```
XX
DΕ
     Synthetic epitope 1 of human cancer antigen eIF3.
XX
KW
     Human; melanoma antigen eukaryotic initiation factor 3; eIF3;
KW
     ovarian cancer; MHC; cytostatic; immunomodulator; immune effector cell;
KW
     anti-cancer; vaccine.
XX
OS
     Homo sapiens.
XX
FH
                     Location/Qualifiers
     Key
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     Domain
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PN
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     06-DEC-2001.
XX
     30-MAY-2001; 2001WO-US017456.
PF
XX
     31-MAY-2000; 2000US-0209391P.
PR
     17-AUG-2000; 2000US-0226258P.
PR
     20-DEC-2000; 2000US-0257008P.
PR
XX
PA
     (GENZ ) GENZYME CORP.
XX
PΙ
     Nicolette CA;
XX
DR
     WPI; 2002-139606/18.
XX
PT
     New therapeutic compounds useful against human ovarian cancer, for
PT
     modulating immune response in a subject, and for generating antibodies
PT
     that specifically recognize and bind to these molecules.
XX
PS
     Claim 29; Page 59; 68pp; English.
XX
     The invention relates to novel therapeutic compounds, that are designed
CC
CC
     to enhance binding to MHC molecules and to enhance immunoregulatory
     properties relative to their natural counterparts. The activity of the
CC
     compounds of the invention may be described as cytostatic and
CC
CC
     immunomodulatory. The compounds are useful against human ovarian cancer,
CC
     for modulating immune response in a subject, and for generating
CC
     antibodies that specifically recognize and bind to these molecules.
CC
     Compositions comprising the compounds are useful as components of anti-
     cancer vaccines and to expand immune effector cells that are specific for
CC
CC
     cells characterised by expression of antigen EIF3 (melanoma antigen
CC
     eukaryotic initiation factor). The peptides or polypeptides conjugated to
CC
     a detectable agent may be used in diagnostic procedures, such as in the
CC
     detection and purification of antibodies, and as immunogens for
CC
     production of antibodies. The polynucleotides can be used as primers for
CC
     detecting genes or gene transcripts expressed in APC to confirm
CC
     transduction of the polynucleotides into host cells. The current sequence
```

Search completed: July 4, 2004, 04:40:35

Job time : 43.8358 secs

## GenCore version 5.1.6 Copyright (c) 1993 - 2004 Compugen Ltd.

OM protein - protein search, using sw model

Run on: July 4, 2004, 04:37:26; Search time 10.6567 Seconds

(without alignments)

82.356 Million cell updates/sec

Title: US-09-641-802-2

Perfect score: 89

Sequence: 1 LQTPQPLLQVMMEPQGD 17

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched: 389414 seqs, 51625971 residues

Total number of hits satisfying chosen parameters: 121837

Minimum DB seq length: 7
Maximum DB seq length: 18

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 100 summaries

Database: Issued Patents AA:\*

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Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

## SUMMARIES

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5	31	34.8	13	2	US-08-780-872-5	Sequence 5, Appli
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8	31	34.8	15	2	US-08-472-281A-30	Sequence 30, Appl
9	31	34.8	15	2	US-08-477-989B-30	Sequence 30, Appl
10	29	32.6	18	2	US-09-017-205-35	Sequence 35, Appl
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## ALIGNMENTS

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RESULT 1
US-09-641-803-2
; Sequence 2, Application US/09641803
 Patent No. 6500798
; GENERAL INFORMATION:
  APPLICANT: STANTON, G. John
  APPLICANT: HUGHES, Thomas K.
; APPLICANT: BOLDOGH, Istvan
  TITLE OF INVENTION: USE OF COLOSTRININ, CONSTITUENT PEPTIDES THEREOF, AND
;
  TITLE OF INVENTION: ANALOGS THEREOF AS OXIDATIVE STRESS REGULATORS
;
   FILE REFERENCE: 265.00220101
   CURRENT APPLICATION NUMBER: US/09/641,803
;
   CURRENT FILING DATE: 2000-08-17
;
   PRIOR APPLICATION NUMBER: 60/149,310
   PRIOR FILING DATE: 1999-08-17
  NUMBER OF SEQ ID NOS: 34
   SOFTWARE: PatentIn Ver. 2.1
; SEQ ID NO 2
   LENGTH: 17
    TYPE: PRT
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ORGANISM: Artificial Sequence
   OTHER INFORMATION: Description of Artificial Sequence: synthetic
   OTHER INFORMATION: peptide
US-09-641-803-2
                        100.0%; Score 89; DB 4; Length 17;
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 Best Local Similarity 100.0%; Pred. No. 7e-08;
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 Matches 17; Conservative
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             Db
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RESULT 2
US-09-641-803-34
; Sequence 34, Application US/09641803
; Patent No. 6500798
; GENERAL INFORMATION:
; APPLICANT: STANTON, G. John
; APPLICANT: HUGHES, Thomas K.
  APPLICANT: BOLDOGH, Istvan
  TITLE OF INVENTION: USE OF COLOSTRININ, CONSTITUENT PEPTIDES THEREOF, AND
  TITLE OF INVENTION: ANALOGS THEREOF AS OXIDATIVE STRESS REGULATORS
  FILE REFERENCE: 265.00220101
; CURRENT APPLICATION NUMBER: US/09/641,803
; CURRENT FILING DATE: 2000-08-17
; PRIOR APPLICATION NUMBER: 60/149,310
; PRIOR FILING DATE: 1999-08-17
; NUMBER OF SEQ ID NOS: 34
  SOFTWARE: PatentIn Ver. 2.1
; SEQ ID NO 34
   LENGTH: 15
   TYPE: PRT
   ORGANISM: Artificial Sequence
   FEATURE:
   OTHER INFORMATION: Description of Artificial Sequence: synthetic
   OTHER INFORMATION: peptide
US-09-641-803-34
  Query Match
                         37.1%; Score 33; DB 4; Length 15;
                        53.8%; Pred. No. 41;
  Best Local Similarity
           7; Conservative 1; Mismatches 5; Indels 0; Gaps
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  Matches
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             1 1111
                     :| |
           3 QPPQPLPPTVMFP 15
Db
RESULT 3
PCT-US91-05177-19
; Sequence 19, Application PC/TUS9105177
; GENERAL INFORMATION:
    APPLICANT: Berkner, Kathy L
    TITLE OF INVENTION: GAMMA-CARBOXYLASE AND METHODS OF USE
    NUMBER OF SEQUENCES: 21
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CORRESPONDENCE ADDRESS:
       ADDRESSEE: Seed and Berry
       STREET: 6300 Columbia Center, 701 Fifth Avenue
      CITY: Seattle
      STATE: WA
;
      COUNTRY: USA
      ZIP: 98104-7092
     COMPUTER READABLE FORM:
      MEDIUM TYPE: Floppy disk
       COMPUTER: IBM PC compatible
      OPERATING SYSTEM: PC-DOS/MS-DOS
       SOFTWARE: PatentIn Release #1.0, Version #1.25
;
     CURRENT APPLICATION DATA:
      APPLICATION NUMBER: PCT/US91/05177
      FILING DATE: 19910722
       CLASSIFICATION: 800
     PRIOR APPLICATION DATA:
      APPLICATION NUMBER: US 07/557,220
;
       FILING DATE: 23-JUL-1990
;
    ATTORNEY/AGENT INFORMATION:
      NAME: Maki, David J
      REGISTRATION NUMBER: 31,392
       REFERENCE/DOCKET NUMBER: 990008.544PC
    TELECOMMUNICATION INFORMATION:
      TELEPHONE: 206-622-4900
       TELEFAX: 206-682-6031
   INFORMATION FOR SEQ ID NO: 19:
     SEQUENCE CHARACTERISTICS:
       LENGTH: 8 amino acids
       TYPE: AMINO ACID
       TOPOLOGY: linear
     FRAGMENT TYPE: internal
PCT-US91-05177-19
  Query Match 36.0%; Score 32; DB 5; Length 8; Best Local Similarity 100.0%; Pred. No. 3e+05;
  Query Match
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  Matches
           6; Conservative
                                                0; Indels
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Qу
              Db
            1 TPQPLL 6
RESULT 4
US-08-162-081B-5
; Sequence 5, Application US/08162081B
; Patent No. 5824492
  GENERAL INFORMATION:
     APPLICANT: Hiles, Ian Donald; Fry, Michael John; Dhand, Ritu
     APPLICANT: Bala; Waterfield, Michael Derek; Parker, Peter
     APPLICANT: Joseph; Otsu, Masayuki; Panayotou, George; Volinia,
     APPLICANT: Stefano; Gout, Ivan Tarasovitch
     TITLE OF INVENTION: POLYPEPTIDES HAVING KINASE ACTIVITY,
     TITLE OF INVENTION: THEIR PREPARATION AND USE
    NUMBER OF SEQUENCES: 50
    CORRESPONDENCE ADDRESS:
      ADDRESSEE: Felfe & Lynch
```

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STREET: 805 Third Avenue
       CITY: New York
       STATE: New York
      COUNTRY: USA
       ZIP: 10022
    COMPUTER READABLE FORM:
      MEDIUM TYPE: Diskette, 5.25 inch, 360 kb storage
      COMPUTER: IBM PS/2
      OPERATING SYSTEM: PC-DOS
       SOFTWARE: Wordperfect
    CURRENT APPLICATION DATA:
      APPLICATION NUMBER: US/08/162,081B
      FILING DATE: February 7, 1994
      CLASSIFICATION: 435
    PRIOR APPLICATION DATA:
      APPLICATION NUMBER: PCT/GB93/00761
      FILING DATE: 13 April 1993
    ATTORNEY/AGENT INFORMATION:
      NAME: Pasqualini, Patricia A.
       REGISTRATION NUMBER: 34,894
       REFERENCE/DOCKET NUMBER: LUD 5256
    TELECOMMUNICATION INFORMATION:
       TELEPHONE: (212) 688-9200
      TELEFAX: (212) 838-3884
  INFORMATION FOR SEQ ID NO: 5:
    SEQUENCE CHARACTERISTICS:
      LENGTH: 13 amino acids
      TYPE: amino acid
      STRANDEDNESS: single
      TOPOLOGY: linear
US-08-162-081B-5
  Query Match
                          34.8%; Score 31; DB 2; Length 13;
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RESULT 5
US-08-780-872-5
; Sequence 5, Application US/08780872
; Patent No. 5846824
  GENERAL INFORMATION:
    APPLICANT: Hiles, Ian Donald; Fry, Michael John; Dhand, Ritu
    APPLICANT: Bala; Waterfield, Michael Derek; Parker, Peter
    APPLICANT: Joseph; Otsu, Masayuki; Panayotou, George; Volinia, APPLICANT: Stefano; Gout, Ivan Tarasovitch
    TITLE OF INVENTION: POLYPEPTIDES HAVING KINASE ACTIVITY,
    TITLE OF INVENTION: THEIR PREPARATION AND USE
    NUMBER OF SEQUENCES: 50
    CORRESPONDENCE ADDRESS:
      ADDRESSEE: Felfe & Lynch
      STREET: 805 Third Avenue
      CITY: New York
```

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STATE: New York
       COUNTRY: USA
       ZIP: 10022
     COMPUTER READABLE FORM:
       MEDIUM TYPE: Diskette, 5.25 inch, 360 kb storage
       COMPUTER: IBM PS/2
       OPERATING SYSTEM: PC-DOS
       SOFTWARE: Wordperfect
     CURRENT APPLICATION DATA:
       APPLICATION NUMBER: US/08/780,872
       FILING DATE: 09-JAN-1997
       CLASSIFICATION: 435
    PRIOR APPLICATION DATA:
      APPLICATION NUMBER: 08/162,081
       FILING DATE: February 7, 1994
       APPLICATION NUMBER: PCT/GB93/00761
       FILING DATE: 13 April 1993
     ATTORNEY/AGENT INFORMATION:
       NAME: Pasqualini, Patricia A.
       REGISTRATION NUMBER: 34,894
       REFERENCE/DOCKET NUMBER: LUD 5256
     TELECOMMUNICATION INFORMATION:
       TELEPHONE: (212) 688-9200
       TELEFAX: (212) 838-3884
   INFORMATION FOR SEQ ID NO: 5:
     SEQUENCE CHARACTERISTICS:
       LENGTH: 13 amino acids
       TYPE: amino acid
       STRANDEDNESS: single
       TOPOLOGY: linear
US-08-780-872-5
  Query Match 34.8%; Score 31; DB 2; Length 13; Best Local Similarity 50.0%; Pred. No. 73;
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Qу
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            4 PDPLYEVMLK 13
RESULT 6
US-09-085-957-5
; Sequence 5, Application US/09085957
; Patent No. 6274327
   GENERAL INFORMATION:
     APPLICANT: Hiles, Ian Donald; Fry, Michael John; Dhand, Ritu
     APPLICANT: Bala; Waterfield, Michael Derek; Parker, Peter
     APPLICANT: Joseph; Otsu, Masayuki; Panayotou, George; Volinia, APPLICANT: Stefano; Gout, Ivan Tarasovitch
     TITLE OF INVENTION: POLYPEPTIDES HAVING KINASE ACTIVITY,
    TITLE OF INVENTION: THEIR PREPARATION AND USE
    NUMBER OF SEQUENCES: 50
    CORRESPONDENCE ADDRESS:
      ADDRESSEE: Felfe & Lynch
       STREET: 805 Third Avenue
       CITY: New York
```

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STATE: New York
      COUNTRY: USA
      ZIP: 10022
    COMPUTER READABLE FORM:
      MEDIUM TYPE: Diskette, 5.25 inch, 360 kb storage
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      COMPUTER: IBM PS/2
      OPERATING SYSTEM: PC-DOS
      SOFTWARE: Wordperfect
    CURRENT APPLICATION DATA:
      APPLICATION NUMBER: US/09/085,957
      FILING DATE:
      CLASSIFICATION:
    PRIOR APPLICATION DATA:
     APPLICATION NUMBER: 08/780,872
      FILING DATE: 09-JAN-1997
      APPLICATION NUMBER: 08/162,081
      FILING DATE: February 7, 1994
      APPLICATION NUMBER: PCT/GB93/00761
      FILING DATE: 13 April 1993
    ATTORNEY/AGENT INFORMATION:
      NAME: Pasqualini, Patricia A.
      REGISTRATION NUMBER: 34,894
      REFERENCE/DOCKET NUMBER: LUD 5256
    TELECOMMUNICATION INFORMATION:
      TELEPHONE: (212) 688-9200
      TELEFAX: (212) 838-3884
  INFORMATION FOR SEQ ID NO: 5:
    SEQUENCE CHARACTERISTICS:
      LENGTH: 13 amino acids
      TYPE: amino acid
      STRANDEDNESS: single
      TOPOLOGY: linear
US-09-085-957-5
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RESULT 7
US-08-477-877B-30
; Sequence 30, Application US/08477877B
; Patent No. 5730979
; GENERAL INFORMATION:
    APPLICANT: Bazin, Herv
    APPLICANT: Latinne, Dominique
    TITLE OF INVENTION: LO-CD2a Antibody and Uses Thereof for Inhibiting T-
Cell Activati
    NUMBER OF SEQUENCES: 96
     CORRESPONDENCE ADDRESS:
      ADDRESSEE: Carella, Byrne, Bain, Gilfillan,
      ADDRESSEE: Cecchi, Stewart & Olstein
      STREET: 6 Becker Farm Road
```

```
CITY: Roseland
      STATE: New Jersey
      COUNTRY: U.S.A.
      ZIP: 07068
    COMPUTER READABLE FORM:
      MEDIUM TYPE: 3.5 inch diskette
      COMPUTER: IBM PS/2
      OPERATING SYSTEM: MS-DOS
      SOFTWARE: WordPerfect 5.1
    CURRENT APPLICATION DATA:
      APPLICATION NUMBER: US/08/477,877B
      FILING DATE: 07-JUN-1995
      CLASSIFICATION: 424
    PRIOR APPLICATION DATA:
      APPLICATION NUMBER: 08/407,009
      FILING DATE: 29-MAR-1995
      APPLICATION NUMBER: 08/119,032
      FILING DATE: 09-SEP-1993
      APPLICATION NUMBER: 08/027,008
      FILING DATE: 05-MAR-1993
    ATTORNEY/AGENT INFORMATION:
     NAME: Olstein, Elliot M.
      REGISTRATION NUMBER: 24,025
      REFERENCE/DOCKET NUMBER: 61750-146
    TELECOMMUNICATION INFORMATION:
      TELEPHONE: 201-994-1700
      TELEFAX: 201-994-1744
  INFORMATION FOR SEQ ID NO: 30:
    SEQUENCE CHARACTERISTICS:
      LENGTH: 15 amino acids
      TYPE: amino acid
      STRANDEDNESS:
      TOPOLOGY: linear
    MOLECULE TYPE: polypeptide
US-08-477-877B-30
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                         34.8%; Score 31; DB 1; Length 15;
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 Matches
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           8 QSPQPLI 14
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RESULT 8
US-08-472-281A-30
; Sequence 30, Application US/08472281A
; Patent No. 5817311
  GENERAL INFORMATION:
    APPLICANT: Bazin, Herv
    APPLICANT: Latinne, Dominique
    TITLE OF INVENTION: LO-CD2a Antibody and Uses Thereof for Inhibiting T-
Cell Activati
    NUMBER OF SEQUENCES: 96
    CORRESPONDENCE ADDRESS:
      ADDRESSEE: Carella, Byrne, Bain, Gilfillan,
```

```
ADDRESSEE: Cecchi, Stewart & Olstein
      STREET: 6 Becker Farm Road
      CITY: Roseland
      STATE: New Jersey
      COUNTRY: U.S.A.
      ZIP: 07068
    COMPUTER READABLE FORM:
      MEDIUM TYPE: 3.5 inch diskette
      COMPUTER: IBM PS/2
      OPERATING SYSTEM: MS-DOS
      SOFTWARE: WordPerfect 5.1
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    CURRENT APPLICATION DATA:
      APPLICATION NUMBER: US/08/472,281A
      FILING DATE: 07-JUN-1995
      CLASSIFICATION: 424
    PRIOR APPLICATION DATA:
      APPLICATION NUMBER: 08/407,009
      FILING DATE: 29-MAR-1995
      APPLICATION NUMBER: 08/119,032
      FILING DATE: 09-SEP-1993
      APPLICATION NUMBER: 08/027,008
      FILING DATE: 05-MAR-1993
    ATTORNEY/AGENT INFORMATION:
      NAME: Olstein, Elliot M.
      REGISTRATION NUMBER: 24,025
      REFERENCE/DOCKET NUMBER: 61750-142
    TELECOMMUNICATION INFORMATION:
      TELEPHONE: 201-994-1700
      TELEFAX: 201-994-1744
  INFORMATION FOR SEQ ID NO:
    SEQUENCE CHARACTERISTICS:
      LENGTH: 15 amino acids
      TYPE: amino acid
      STRANDEDNESS:
      TOPOLOGY: linear
    MOLECULE TYPE: polypeptide
US-08-472-281A-30
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  Best Local Similarity 71.4%; Pred. No. 85;
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 Matches
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Qу
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Db
           8 QSPQPLI 14
RESULT 9
US-08-477-989B-30
; Sequence 30, Application US/08477989B
; Patent No. 5951983
  GENERAL INFORMATION:
    APPLICANT: Bazin, Herv
    APPLICANT: Latinne, Dominique
    APPLICANT: Kaplan, Ruth
    APPLICANT: Kieber-Emmons, Thomas
    APPLICANT: Postema, Christina E.
```

```
APPLICANT: White-Scharf, Mary
    TITLE OF INVENTION: LO-CD2a Antibody and Uses
    TITLE OF INVENTION: Thereof for Inhibiting
    TITLE OF INVENTION: T-Cell Activation and
    TITLE OF INVENTION: Proliferation
    NUMBER OF SEQUENCES: 96
    CORRESPONDENCE ADDRESS:
      ADDRESSEE: Carella, Byrne, Bain, Gilfillan,
      ADDRESSEE: Cecchi, Stewart & Olstein
      STREET: 6 Becker Farm Road
      CITY: Roseland
      STATE: New Jersey
      COUNTRY: U.S.A.
      ZIP: 07068
    COMPUTER READABLE FORM:
      MEDIUM TYPE: 3.5 inch diskette
      COMPUTER: IBM PS/2
      OPERATING SYSTEM: MS-DOS
      SOFTWARE: WordPerfect 5.1
    CURRENT APPLICATION DATA:
      APPLICATION NUMBER: US/08/477,989B
      FILING DATE: 07-JUN-1995
      CLASSIFICATION: 424
    PRIOR APPLICATION DATA:
     APPLICATION NUMBER: 08/407,009
      FILING DATE: 29-MAR-1995
      APPLICATION NUMBER: 08/119,032
      FILING DATE: 09-SEP-1993
      APPLICATION NUMBER: 08/027,008
      FILING DATE: 05-MAR-1993
    ATTORNEY/AGENT INFORMATION:
      NAME: Olstein, Elliot M.
      REGISTRATION NUMBER: 24,025
      REFERENCE/DOCKET NUMBER: 61750-147
    TELECOMMUNICATION INFORMATION:
      TELEPHONE: 201-994-1700
      TELEFAX: 201-994-1744
  INFORMATION FOR SEQ ID NO: 30:
    SEQUENCE CHARACTERISTICS:
      LENGTH: 15 amino acids
      TYPE: amino acid
      STRANDEDNESS:
      TOPOLOGY: linear
    MOLECULE TYPE: polypeptide
US-08-477-989B-30
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 Best Local Similarity 71.4%; Pred. No. 85;
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RESULT 10 US-09-017-205-35

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; Sequence 35, Application US/09017205
; Patent No. 5965357
  GENERAL INFORMATION:
    APPLICANT: Marsden, Howard S
    TITLE OF INVENTION: PEPTIDE STRUCTURES AND THEIR USE IN
    TITLE OF INVENTION: DIAGNOSIS OF HERPES SIMPLEX VIRUS TYPE 2
    NUMBER OF SEQUENCES: 86
    CORRESPONDENCE ADDRESS:
      ADDRESSEE: Nixon & Vanderhye PC
       STREET: 8th Floor, 1100 No. 5965357th Glebe Road
      CITY: Arlington
      STATE: Virginia
      COUNTRY: USA
       ZIP: 22201-4714
    COMPUTER READABLE FORM:
      MEDIUM TYPE: Floppy disk
      COMPUTER: IBM PC compatible
       OPERATING SYSTEM: PC-DOS/MS-DOS
       SOFTWARE: PatentIn Release #1.0, Version #1.30
    CURRENT APPLICATION DATA:
      APPLICATION NUMBER: US/09/017,205
      FILING DATE: 02-FEB-1998
      CLASSIFICATION: 435
    ATTORNEY/AGENT INFORMATION:
      NAME: Mitchard, Leonard C
      REGISTRATION NUMBER: 29,009
      REFERENCE/DOCKET NUMBER: 604-436
    TELECOMMUNICATION INFORMATION:
      TELEPHONE: (703)816-4000
       TELEFAX: (703)816-4100
   INFORMATION FOR SEQ ID NO: 35:
    SEQUENCE CHARACTERISTICS:
       LENGTH: 18 amino acids
       TYPE: amino acid
       TOPOLOGY: linear
    MOLECULE TYPE: peptide from HSV-2 glycoprotein G
     FRAGMENT TYPE: internal
US-09-017-205-35
  Query Match
                         32.6%; Score 29; DB 2; Length 18;
                         37.5%; Pred. No. 2.1e+02;
  Best Local Similarity
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                                2; Mismatches
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RESULT 11
US-08-594-447-74
; Sequence 74, Application US/08594447
; Patent No. 5776716
  GENERAL INFORMATION:
    APPLICANT: Ron, Dorit
    APPLICANT: Napolitano, Eugene W.
    APPLICANT: Voronova, Anna F.
    TITLE OF INVENTION: METHODS FOR IDENTIFYING AGENTS WHICH
```

```
TITLE OF INVENTION: BLOCKTHE INTERACTION OF FYN WITH PKC-THETA, AND USES
    TITLE OF INVENTION: THEREOF
    NUMBER OF SEQUENCES: 75
    CORRESPONDENCE ADDRESS:
      ADDRESSEE: MORRISON & FOERSTER
      STREET: 2000 Pennsylvania Avenue, NW - Ste. 5500
      CITY: Washington
      STATE: DC
      COUNTRY: USA
      ZIP: 20006-1888
    COMPUTER READABLE FORM:
      MEDIUM TYPE: Floppy disk
      COMPUTER: IBM PC compatible
      OPERATING SYSTEM: PC-DOS/MS-DOS
      SOFTWARE: PatentIn Release #1.0, Version #1.30
    CURRENT APPLICATION DATA:
      APPLICATION NUMBER: US/08/594,447
      FILING DATE: 31-JAN-1996
      CLASSIFICATION: 435
    ATTORNEY/AGENT INFORMATION:
      NAME: Murashige, Kate H.
      REGISTRATION NUMBER: 29,959
      REFERENCE/DOCKET NUMBER: 22550-20025.24
    TELECOMMUNICATION INFORMATION:
      TELEPHONE: (202) 887-1500
      TELEFAX: (202) 822-0168
      TELEX: 90-4030 MRSNFOERSWSH
  INFORMATION FOR SEQ ID NO: 74:
    SEQUENCE CHARACTERISTICS:
      LENGTH: 7 amino acids
      TYPE: amino acid
      STRANDEDNESS: single
      TOPOLOGY: linear
    MOLECULE TYPE: peptide
    FEATURE:
      NAME/KEY: Peptide
      LOCATION: 1..7
      OTHER INFORMATION: /label= PRK1-3
US-08-594-447-74
 Query Match
                         31.5%; Score 28; DB 1; Length 7;
                         100.0%; Pred. No. 3e+05;
  Best Local Similarity
            5; Conservative 0; Mismatches 0; Indels
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          12 MEPQG 16
Qу
             +11111
           1 MEPQG 5
RESULT 12
US-08-541-964-73
; Sequence 73, Application US/08541964
; Patent No. 5783405
  GENERAL INFORMATION:
    APPLICANT: Mochly-Rosen, Daria
    APPLICANT: Ron, Dorit
    APPLICANT: Kauvar, Lawrence M.
```

```
APPLICANT: Napolitano, Eugene W.
    TITLE OF INVENTION: A RAPID SCREENING METHOD FOR EFFECTORS
    TITLE OF INVENTION: OF SIGNAL TRANSDUCTION
    NUMBER OF SEQUENCES: 74
    CORRESPONDENCE ADDRESS:
      ADDRESSEE: MORRISON & FOERSTER
      STREET: 2000 PENNSYLVANIA AVENUE, NW-STE. 5500
      CITY: WASHINGTON
      STATE: DC
      COUNTRY: USA
      ZIP: 20006-1888
    COMPUTER READABLE FORM:
      MEDIUM TYPE: Floppy disk
      COMPUTER: IBM PC compatible
      OPERATING SYSTEM: PC-DOS/MS-DOS
      SOFTWARE: PatentIn Release #1.0, Version #1.30
    CURRENT APPLICATION DATA:
      APPLICATION NUMBER: US/08/541,964
      FILING DATE: 10-OCT-1995
      CLASSIFICATION: 435
    ATTORNEY/AGENT INFORMATION:
      NAME: Murashige, Kate H.
      REGISTRATION NUMBER: 29,959
      REFERENCE/DOCKET NUMBER: 22550-20025.23
    TELECOMMUNICATION INFORMATION:
      TELEPHONE: (202) 887-1500
      TELEFAX: (202) 822-0168
      TELEX: 90-4030 MRSNFOERSWSH
  INFORMATION FOR SEQ ID NO: 73:
    SEOUENCE CHARACTERISTICS:
      LENGTH: 7 amino acids
      TYPE: amino acid
      STRANDEDNESS: single
      TOPOLOGY: linear
    MOLECULE TYPE: peptide
    FEATURE:
      NAME/KEY: Peptide
      LOCATION: 1...7
      OTHER INFORMATION: /label= PRK1-3
US-08-541-964-73
                         31.5%; Score 28; DB 1; Length 7;
 Query Match
                        100.0%; Pred. No. 3e+05;
 Best Local Similarity
 Matches 5; Conservative 0; Mismatches 0; Indels 0; Gaps
                                                                           0;
          12 MEPQG 16
Qу
             \Box
           1 MEPQG 5
Db
RESULT 13
US-08-665-647-88
; Sequence 88, Application US/08665647
; Patent No. 5935803
; GENERAL INFORMATION:
    APPLICANT: Dasquez, Nicki J.
    APPLICANT: Ron, Dorit
```

```
APPLICANT: Voronova, Anna F.
    APPLICANT: Napolitano, Eugene W.
    TITLE OF INVENTION: METHODS TO IDENTIFY IMMUNOMODULATORS
    TITLE OF INVENTION: USING COGNATE INTERACTION OF PKC-THETA
    NUMBER OF SEQUENCES: 89
;
    CORRESPONDENCE ADDRESS:
      ADDRESSEE: MORRISON & FOERSTER
      STREET: 2000 Pennsylvania Avenue, NW - Ste. 5500
      CITY: Washington
      STATE: DC
      COUNTRY: USA
      ZIP: 20006-1888
    COMPUTER READABLE FORM:
      MEDIUM TYPE: Floppy disk
      COMPUTER: IBM PC compatible
      OPERATING SYSTEM: PC-DOS/MS-DOS
      SOFTWARE: PatentIn Release #1.0, Version #1.30
    CURRENT APPLICATION DATA:
      APPLICATION NUMBER: US/08/665,647
;
      FILING DATE: 18-JUN-1996
;
      CLASSIFICATION: 435
    ATTORNEY/AGENT INFORMATION:
      NAME: Murashige, Kate H.
;
      REGISTRATION NUMBER: 29,959
      REFERENCE/DOCKET NUMBER: 22550-20025.25
    TELECOMMUNICATION INFORMATION:
      TELEPHONE: (202) 887-1500
;
      TELEFAX: (202) 822-0168
      TELEX: 90-4030 MRSNFOERSWSH
  INFORMATION FOR SEQ ID NO: 88:
    SEQUENCE CHARACTERISTICS:
      LENGTH: 7 amino acids
      TYPE: amino acid
;
      STRANDEDNESS: single
      TOPOLOGY: linear
;
    MOLECULE TYPE: peptide
    FEATURE:
      NAME/KEY: Peptide
      LOCATION: 1..7
      OTHER INFORMATION: /label= PRK1-3
US-08-665-647-88
                         31.5%; Score 28; DB 2; Length 7;
 Query Match
 Best Local Similarity 100.0%; Pred. No. 3e+05;
 Matches 5; Conservative 0; Mismatches 0; Indels
                                                               0; Gaps
          12 MEPQG 16
Qу
             11111
Db
           1 MEPQG 5
RESULT 14
US-08-981-392-53
; Sequence 53, Application US/08981392
; Patent No. 6262025
; GENERAL INFORMATION:
    APPLICANT: Ish-Horowicz, David
```

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APPLICANT: Henrique, Domingos Manuel Pinto
    APPLICANT: Lewis, Julian Hart
    APPLICANT: Artavanis-Tsakonas, Spyridon
    APPLICANT: Gray, Grace
    TITLE OF INVENTION: NUCLEOTIDE AND PROTEIN SEQUENCES
    TITLE OF INVENTION: OF VERTEBRATE DELTA GENES AND METHODS BASED THEREON
    NUMBER OF SEQUENCES: 94
    CORRESPONDENCE ADDRESS:
      ADDRESSEE: Pennie & Edmonds LLP
      STREET: 1155 Avenue of the Americas
      CITY: New York
      STATE: NY
      COUNTRY: USA
      ZIP: 10036/2711
    COMPUTER READABLE FORM:
      MEDIUM TYPE: Diskette
      COMPUTER: IBM Compatible
      OPERATING SYSTEM: DOS
      SOFTWARE: FastSEQ Version 2.0
    CURRENT APPLICATION DATA:
      APPLICATION NUMBER: US/08/981,392
      FILING DATE: 22-DEC-1997
      CLASSIFICATION: 514
    ATTORNEY/AGENT INFORMATION:
      NAME: Antler, Adriane M.
      REGISTRATION NUMBER: 32,605
      REFERENCE/DOCKET NUMBER: 7326-038
    TELECOMMUNICATION INFORMATION:
      TELEPHONE: 212-790-9090
      TELEFAX: 212-869-8864
      TELEX: 66141 PENNIE
  INFORMATION FOR SEO ID NO: 53:
    SEQUENCE CHARACTERISTICS:
      LENGTH: 10 amino acids
      TYPE: amino acid
      STRANDEDNESS:
      TOPOLOGY: unknown
    MOLECULE TYPE: peptide
US-08-981-392-53
 Query Match
                         31.5%; Score 28; DB 3; Length 10;
 Best Local Similarity 50.0%; Pred. No. 1.6e+02;
           5; Conservative 2; Mismatches 3; Indels
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                                                                           0;
Qу
           4 PQPLLQVMME 13
             1111::
           1 PQPLVRTEQE 10
RESULT 15
US-08-553-257A-21
; Sequence 21, Application US/08553257A
; Patent No. 5994083
  GENERAL INFORMATION:
    APPLICANT: ISTITUTO DI RICERCHE DI BIOLOGIA
    APPLICANT: MOLECOLARE P. ANGELETTI S.p.A.
    APPLICANT: FELICI, Franco
```

```
APPLICANT: LUZZAGO, Alessandra
    APPLICANT: NICOSIA, Alfredo
    APPLICANT: MONACI, Paolo
    APPLICANT: CORTESE, Riccardo
    TITLE OF INVENTION: PROCESS FOR THE PREPARATION OF IMMUNOGENS TITLE OF INVENTION: OR DIAGNOSTIC REAGENTS, AND IMMUNOGENS OR
;
    TITLE OF INVENTION: DIAGNOSTIC REAGENTS THEREBY OBTAINABLE
    NUMBER OF SEQUENCES: 68
    CORRESPONDENCE ADDRESS:
      ADDRESSEE: Browdy and Neimark
      STREET: 419 Seventh Street N.W. Ste. 300
      CITY: Washington
;
      STATE: D.C.
      COUNTRY: USA
      ZIP: 20004
    COMPUTER READABLE FORM:
      MEDIUM TYPE: Floppy disk
      COMPUTER: IBM PC compatible
      OPERATING SYSTEM: PC-DOS/MS-DOS
;
      SOFTWARE: PatentIn Release #1.0, Version #1.30
    CURRENT APPLICATION DATA:
     APPLICATION NUMBER: US/08/553,257A
     FILING DATE:
;
      CLASSIFICATION: 435
    PRIOR APPLICATION DATA:
      APPLICATION NUMBER: PCT/IT94/00054
      FILING DATE: 05-MAY-1994
    PRIOR APPLICATION DATA:
     APPLICATION NUMBER: RM93A000301
      FILING DATE: 11-MAY-1993
    ATTORNEY/AGENT INFORMATION:
      NAME: Browdy, Roger L.
      REGISTRATION NUMBER: 25,618
      REFERENCE/DOCKET NUMBER: FELICI=1
    TELECOMMUNICATION INFORMATION:
      TELEPHONE: (202) 628-5197
       TELEFAX: (202) 737-3528
   INFORMATION FOR SEQ ID NO: 21:
    SEQUENCE CHARACTERISTICS:
      LENGTH: 15 amino acids
      TYPE: amino acid
;
       STRANDEDNESS: single
      TOPOLOGY: linear
   MOLECULE TYPE: recombinant protein
   HYPOTHETICAL: yes
    FRAGMENT TYPE: internal
    IMMEDIATE SOURCE:
      LIBRARY: of recombinant peptides on phage
       CLONE: phagic
     FEATURE:
       NAME/KEY: polypeptide
       IDENTIFICATION METHOD: selection with specific antibodies
US-08-553-257A-21
  Query Match
                          31.5%; Score 28; DB 2; Length 15;
  Best Local Similarity 50.0%; Pred. No. 2.5e+02;
  Matches 8; Conservative 0; Mismatches 4; Indels 4; Gaps
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2 QTPQPLLQVMMEPQGD 17
Qу
              1 11 1
                         11 11
Db
            4 QVPQSRL----EPWGD 15
RESULT 16
US-09-441-992-21
; Sequence 21, Application US/09441992
; Patent No. 6541210
   GENERAL INFORMATION:
         APPLICANT: ISTITUTO DI RICERCHE DI BIOLOGIA
                    MOLECOLARE P. ANGELETTI S.p.A.
                    FELICI, Franco
                    LUZZAGO, Alessandra
                    NICOSIA, Alfredo
                    MONACI, Paolo
                    CORTESE, Riccardo
         TITLE OF INVENTION: PROCESS FOR THE PREPARATION OF IMMUNOGENS
                             OR DIAGNOSTIC REAGENTS, AND IMMUNOGENS OR
                             DIAGNOSTIC REAGENTS THEREBY OBTAINABLE
        NUMBER OF SEQUENCES: 68
         CORRESPONDENCE ADDRESS:
              ADDRESSEE: Browdy and Neimark
              STREET: 419 Seventh Street N.W. Ste. 300
              CITY: Washington
              STATE: D.C.
              COUNTRY: USA
              ZIP: 20004
         COMPUTER READABLE FORM:
              MEDIUM TYPE: Floppy disk
              COMPUTER: IBM PC compatible
              OPERATING SYSTEM: PC-DOS/MS-DOS
              SOFTWARE: PatentIn Release #1.0, Version #1.30
         CURRENT APPLICATION DATA:
              APPLICATION NUMBER: US/09/441,992
              FILING DATE: 18-No. 6541210-1999
              CLASSIFICATION: <Unknown>
         PRIOR APPLICATION DATA:
              APPLICATION NUMBER: 08/553,257
              FILING DATE: <Unknown>
              APPLICATION NUMBER: RM93A000301
              FILING DATE: 11-MAY-1993
        ATTORNEY/AGENT INFORMATION:
              NAME: Browdy, Roger L.
              REGISTRATION NUMBER: 25,618
              REFERENCE/DOCKET NUMBER: FELICI=1
         TELECOMMUNICATION INFORMATION:
              TELEPHONE: (202) 628-5197
              TELEFAX: (202) 737-3528
    INFORMATION FOR SEQ ID NO: 21:
         SEQUENCE CHARACTERISTICS:
              LENGTH: 15 amino acids
              TYPE: amino acid
              STRANDEDNESS: single
              TOPOLOGY: linear
         MOLECULE TYPE: recombinant protein
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HYPOTHETICAL: yes
         FRAGMENT TYPE: internal
         IMMEDIATE SOURCE:
              LIBRARY: of recombinant peptides on phage
              CLONE: phagic
                     (ix)
                          FEATURE
                     (A) NAME: polypeptide
         SEQUENCE DESCRIPTION: SEQ ID NO: 21:
US-09-441-992-21
                          31.5%; Score 28; DB 4; Length 15;
  Query Match
  Best Local Similarity 50.0%; Pred. No. 2.5e+02;
            8; Conservative 0; Mismatches 4; Indels
                                                                  4; Gaps
                                                                              1;
Qу
            2 QTPQPLLQVMMEPQGD 17
             11 11
Db
            4 QVPQSRL----EPWGD 15
RESULT 17
US-08-602-999A-195
; Sequence 195, Application US/08602999A
; Patent No. 6184205
  GENERAL INFORMATION:
    APPLICANT: SPARKS, Andrew B.
    APPLICANT: KAY, Brian K.
APPLICANT: THORN, Judith M.
APPLICANT: QUILLIAM, Lawrence A.
    APPLICANT: DER, Channing J.
    APPLICANT: FOWLKES, Dana M.
    APPLICANT: RIDER, James E.
    TITLE OF INVENTION: SH3 BINDING PEPTIDES AND METHODS OF
     TITLE OF INVENTION: ISOLATING AND USING SAME
    NUMBER OF SEQUENCES: 467
    CORRESPONDENCE ADDRESS:
       ADDRESSEE: Pennie & Edmonds
       STREET: 1155 Avenue of the Americas
      CITY: New York
       STATE: New York
       COUNTRY: U.S.A.
       ZIP: 10036-2711
    COMPUTER READABLE FORM:
       MEDIUM TYPE: Floppy disk
       COMPUTER: IBM PC compatible
       OPERATING SYSTEM: PC-DOS/MS-DOS
       SOFTWARE: PatentIn Release #1.0, Version #1.30
     CURRENT APPLICATION DATA:
       APPLICATION NUMBER: US/08/602,999A
       FILING DATE: 16-FEB-1996
       CLASSIFICATION: 435
     ATTORNEY/AGENT INFORMATION:
       NAME: Misrock, S. Leslie
       REGISTRATION NUMBER: 18,872
       REFERENCE/DOCKET NUMBER: 1101-202
;
     TELECOMMUNICATION INFORMATION:
       TELEPHONE: (212) 790-9090
       TELEFAX: (212) 869-9741/8864
```

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TELEX: 66141 PENNIE
   INFORMATION FOR SEQ ID NO: 195:
     SEQUENCE CHARACTERISTICS:
       LENGTH: 16 amino acids
       TYPE: amino acid
       TOPOLOGY: unknown
     MOLECULE TYPE: peptide
US-08-602-999A-195
                          31.5%; Score 28; DB 3; Length 16;
  Ouerv Match
  Best Local Similarity 41.7%; Pred. No. 2.7e+02;
           5; Conservative 2; Mismatches 5; Indels
  Matches
                                                                0; Gaps
            3 TPQPLLQVMMEP 14
Qу
             11:1: 11
Db
            4 TPRPAVPQRMNP 15
RESULT 18
US-09-500-124-195
; Sequence 195, Application US/09500124
; Patent No. 6432920
   GENERAL INFORMATION:
    APPLICANT: SPARKS, Andrew B.
    APPLICANT: KAY, Brian K.
    APPLICANT: THORN, Judith M.
    APPLICANT: QUILLIAM, Lawrence A. APPLICANT: DER, Channing J.
    APPLICANT: FOWLKES, Dana M.
    APPLICANT: RIDER, James E.
    TITLE OF INVENTION: SH3 BINDING PEPTIDES AND METHODS OF
    TITLE OF INVENTION: ISOLATING AND USING SAME
    NUMBER OF SEQUENCES: 467
    CORRESPONDENCE ADDRESS:
      ADDRESSEE: Pennie & Edmonds
       STREET: 1155 Avenue of the Americas
      CITY: New York
       STATE: New York
      COUNTRY: U.S.A.
       ZIP: 10036-2711
    COMPUTER READABLE FORM:
      MEDIUM TYPE: Floppy disk
       COMPUTER: IBM PC compatible
       OPERATING SYSTEM: PC-DOS/MS-DOS
       SOFTWARE: PatentIn Release #1.0, Version #1.30
     CURRENT APPLICATION DATA:
       APPLICATION NUMBER: US/09/500,124
      FILING DATE:
       CLASSIFICATION:
     PRIOR APPLICATION DATA:
       APPLICATION NUMBER: 08/602,999
       FILING DATE: 16-FEB-1996
     ATTORNEY/AGENT INFORMATION:
       NAME: Misrock, S. Leslie
       REGISTRATION NUMBER: 18,872
       REFERENCE/DOCKET NUMBER: 1101-202
     TELECOMMUNICATION INFORMATION:
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```
TELEPHONE: (212) 790-9090
       TELEFAX: (212) 869-9741/8864
       TELEX: 66141 PENNIE
  INFORMATION FOR SEQ ID NO: 195:
    SEQUENCE CHARACTERISTICS:
       LENGTH: 16 amino acids
       TYPE: amino acid
       TOPOLOGY: unknown
    MOLECULE TYPE: peptide
US-09-500-124-195
                         31.5%; Score 28; DB 4; Length 16;
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 Best Local Similarity 41.7%; Pred. No. 2.7e+02;
  Matches
           5; Conservative 2; Mismatches
                                               5; Indels 0; Gaps
                                                                           0;
Qy
           3 TPQPLLQVMMEP 14
             11:1: 11
Db
           4 TPRPAVPQRMNP 15
RESULT 19
US-08-646-265A-125
; Sequence 125, Application US/08646265A
; Patent No. 6214973
  GENERAL INFORMATION:
    APPLICANT: OHTOMO, Toshihiko
    APPLICANT: SATO, Koh
    APPLICANT: TSUCHIYA, Masayuki
    TITLE OF INVENTION: RESHAPED HUMAN ANTIBODY TO HUMAN
    TITLE OF INVENTION: MEDULLOBLASTOMA CELLS
    NUMBER OF SEQUENCES: 132
    CORRESPONDENCE ADDRESS:
      ADDRESSEE: Foley & Lardner
       STREET: 3000 K Street, N.W., Suite 500
      CITY: Washington
       STATE: D.C.
;
      COUNTRY: USA
       ZIP: 20007-5109
    COMPUTER READABLE FORM:
      MEDIUM TYPE: Floppy disk
      COMPUTER: IBM PC compatible
      OPERATING SYSTEM: PC-DOS/MS-DOS
       SOFTWARE: PatentIn Release #1.0, Version #1.30
    CURRENT APPLICATION DATA:
      APPLICATION NUMBER: US/08/646,265A
       FILING DATE: 09-SEP-1996
       CLASSIFICATION: 435
     PRIOR APPLICATION DATA:
      APPLICATION NUMBER: WO PCT/JP94/01763
       FILING DATE: 19-OCT-1994
     PRIOR APPLICATION DATA:
       APPLICATION NUMBER: JP 5-291078
       FILING DATE: 19-NOV-1993
    ATTORNEY/AGENT INFORMATION:
      NAME: WEGNER, Harold C.
       REGISTRATION NUMBER: 25,258
       REFERENCE/DOCKET NUMBER: 53466/184
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TELECOMMUNICATION INFORMATION:
       TELEPHONE: (202) 672-5300
      TELEFAX: (202) 672-5399
       TELEX: 904136
   INFORMATION FOR SEQ ID NO: 125:
    SEQUENCE CHARACTERISTICS:
      LENGTH: 15 amino acids
      TYPE: amino acid
      TOPOLOGY: linear
    MOLECULE TYPE: protein
US-08-646-265A-125
  Query Match
                         30.3%; Score 27; DB 3; Length 15;
  Best Local Similarity 57.1%; Pred. No. 3.6e+02;
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          4; Conservative 3; Mismatches 0; Indels
                                                               0; Gaps
                                                                           0;
Qу
           2 QTPQPLL 8
             1:1:11:
Db
           8 QSPKPLI 14
RESULT 20
US-09-947-372A-27
; Sequence 27, Application US/09947372A
; Patent No. 6613557
; GENERAL INFORMATION:
 APPLICANT: FRAZER, IAN
  APPLICANT: ZHOU, JIAN
  TITLE OF INVENTION: PAPILLOMAVIRUS VACCINE
; FILE REFERENCE: 065064/0137
  CURRENT APPLICATION NUMBER: US/09/947,372A
  CURRENT FILING DATE: 2001-09-07
; PRIOR APPLICATION NUMBER: 08/185,928
  PRIOR FILING DATE: 1994-01-19
  PRIOR APPLICATION NUMBER: PCT/AU92/02184
  PRIOR FILING DATE: 1992-07-20
  PRIOR APPLICATION NUMBER: AU PK7322
  PRIOR FILING DATE: 1991-07-19
; NUMBER OF SEQ ID NOS: 66
  SOFTWARE: PatentIn Ver. 2.1
; SEQ ID NO 27
   LENGTH: 15
   TYPE: PRT
   ORGANISM: Human papillomavirus type 16
US-09-947-372A-27
                         30.3%; Score 27; DB 4; Length 15;
  Query Match
  Best Local Similarity 40.0%; Pred. No. 3.6e+02;
                             4; Mismatches 2; Indels
 Matches
           4; Conservative
                                                               0; Gaps
                                                                           0;
           8 LQVMMEPQGD 17
Qу
             :::: || ||
           1 IKMVSEPYGD 10
Db
RESULT 21
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RESULT 21 US-08-837-226-2

```
; Sequence 2, Application US/08837226
; Patent No. 6043216
; GENERAL INFORMATION:
     APPLICANT: Toback, F. Gary
     APPLICANT: Lieske, John C.
     TITLE OF INVENTION: METHODS AND COMPOSITION FOR DETECTING
TITLE OF INVENTION: AND TREATING KIDNEY DISEASES ASSOCIATED WITH ADHESION
OF
    TITLE OF INVENTION: CRYSTALS TO KIDNEY CELLS
    NUMBER OF SEQUENCES: 14
    CORRESPONDENCE ADDRESS:
     ADDRESSEE: BRINKS, HOFER, GILSON & LIONE
      STREET: NBC Tower - Suite 3600, 455 N. Cityfront
      STREET: Plaza Drive
      CITY: Chicago
      STATE: Illinois
      COUNTRY: USA
      ZIP: 60611-5599
    COMPUTER READABLE FORM:
      MEDIUM TYPE: Floppy disk
      COMPUTER: IBM PC compatible
      OPERATING SYSTEM: PC-DOS/MS-DOS
       SOFTWARE: PatentIn Release #1.0, Version #1.30
    CURRENT APPLICATION DATA:
      APPLICATION NUMBER: US/08/837,226
      FILING DATE: 08-APR-1997
      CLASSIFICATION: 530
    PRIOR APPLICATION DATA:
     APPLICATION NUMBER: US 08/389,005
      FILING DATE: 15-FEB-1995
    ATTORNEY/AGENT INFORMATION:
      NAME: Martin, Alice O.
      REGISTRATION NUMBER: 35,601
       REFERENCE/DOCKET NUMBER: 7814/24
     TELECOMMUNICATION INFORMATION:
       TELEPHONE: 312-321-4200
       TELEFAX: 312-321-4299
   INFORMATION FOR SEQ ID NO: 2:
    SEQUENCE CHARACTERISTICS:
      LENGTH: 16 amino acids
      TYPE: amino acid
      STRANDEDNESS:
      TOPOLOGY: linear
    MOLECULE TYPE: peptide
US-08-837-226-2
                          30.3%; Score 27; DB 3; Length 16;
  Query Match
  Best Local Similarity 83.3%; Pred. No. 3.9e+02;
  Matches 5; Conservative 0; Mismatches 1; Indels 0; Gaps
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Qy
            4 PQPLLQ 9
             111111
Db
           10 PQPLYQ 15
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RESULT 22 US-09-537-226-2

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; Sequence 2, Application US/09537226
; Patent No. 6482934
; GENERAL INFORMATION:
  APPLICANT: TOBACK, F. GARY
  APPLICANT: LIESKE, JOHN C.
  TITLE OF INVENTION: METHODS AND COMPOSITIONS FOR DETECTING AND TREATING
  TITLE OF INVENTION: KIDNEY DISEASES ASSOCIATED WITH ADHESION OF CRYSTALS TO
  TITLE OF INVENTION: KIDNEY CELLS
  FILE REFERENCE: 21459/90606
  CURRENT APPLICATION NUMBER: US/09/537,226
  CURRENT FILING DATE: 2000-03-28
  PRIOR APPLICATION NUMBER: 08/389,005
  PRIOR FILING DATE: 1995-02-15
  NUMBER OF SEQ ID NOS: 23
  SOFTWARE: PatentIn Ver. 2.1
; SEQ ID NO 2
   LENGTH: 16
   TYPE: PRT
   ORGANISM: Artificial Sequence
   FEATURE:
   OTHER INFORMATION: Description of Artificial Sequence: Synthetic
   OTHER INFORMATION: peptide
US-09-537-226-2
  Query Match
                         30.3%; Score 27; DB 4; Length 16;
 Best Local Similarity 83.3%; Pred. No. 3.9e+02;
           5; Conservative 0; Mismatches 1; Indels
 Matches
                                                                0; Gaps
                                                                            0;
           4 PQPLLQ 9
Qу
             ++++
Db
          10 PQPLYQ 15
RESULT 23
US-09-829-855-240
; Sequence 240, Application US/09829855
; Patent No. 6613520
; GENERAL INFORMATION:
; APPLICANT: Matthew, Ashby N.
  TITLE OF INVENTION: Methods for the Survey and Genetic Analysis of
Populations
 FILE REFERENCE: ASHBY-1
  CURRENT APPLICATION NUMBER: US/09/829,855
  CURRENT FILING DATE: 2001-04-10
  PRIOR APPLICATION NUMBER: US 60/196063
  PRIOR FILING DATE: 2000-04-10
  PRIOR APPLICATION NUMBER: US 60/196258
  PRIOR FILING DATE: 2000-04-11
; NUMBER OF SEQ ID NOS: 244
  SOFTWARE: PatentIn version 3.1
; SEQ ID NO 240
   LENGTH: 18
   TYPE: PRT
   ORGANISM: Unknown
   FEATURE:
   OTHER INFORMATION: Uncultured Acidobacterium Sub.Div-2
US-09-829-855-240
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Query Match
                       29.8%; Score 26.5; DB 4; Length 18;
  Best Local Similarity 58.3%; Pred. No. 5.3e+02;
  Matches 7; Conservative 1; Mismatches 3; Indels 1; Gaps
                                                                         1;
           5 QPLLQVMMEPQG 16
Qу
             11 | 1: 111
Db
           3 QP-LHVVATPQG 13
RESULT 24
US-08-475-955-182
; Sequence 182, Application US/08475955
; Patent No. 6641813
  GENERAL INFORMATION:
    APPLICANT: Harley, John
    TITLE OF INVENTION: METHODS AND REAGENTS FOR DIAGNOSIS OF
    TITLE OF INVENTION: AUTOANTIBODIES
    NUMBER OF SEQUENCES: 218
   CORRESPONDENCE ADDRESS:
     ADDRESSEE: Patrea L. Pabst
      STREET: 2800 One Atlantic Center, 1250 West Peachtree Street
      CITY: Atlanta
     STATE: GA
      COUNTRY: USA
      ZIP: 30309-3450
    COMPUTER READABLE FORM:
      MEDIUM TYPE: Floppy disk
      COMPUTER: IBM PC compatible
      OPERATING SYSTEM: PC-DOS/MS-DOS
      SOFTWARE: PatentIn Release #1.0, Version #1.25
    CURRENT APPLICATION DATA:
      APPLICATION NUMBER: US/08/475,955
      FILING DATE: June 7, 1995
      CLASSIFICATION: 424
    PRIOR APPLICATION DATA:
      APPLICATION NUMBER: 07/867,819
      FILING DATE: April 13, 1992
      CLASSIFICATION: 424
    PRIOR APPLICATION DATA:
      APPLICATION NUMBER: 07/648,205
      FILING DATE: January 31, 1991
    PRIOR APPLICATION DATA:
      APPLICATION NUMBER: 07/472,947
      FILING DATE: January 31, 1990
    ATTORNEY/AGENT INFORMATION:
      NAME: Pabst, Patrea L.
      REGISTRATION NUMBER: 31,284
      REFERENCE/DOCKET NUMBER: OMRF114CIP(2) DIV
    TELECOMMUNICATION INFORMATION:
      TELEPHONE: (404)-873-8794
       TELEFAX: (404)-873-8795
   INFORMATION FOR SEQ ID NO: 182:
     SEQUENCE CHARACTERISTICS:
      LENGTH: 8 amino acids
      TYPE: amino acid
;
      STRANDEDNESS: single
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TOPOLOGY: linear
    MOLECULE TYPE: peptide
US-08-475-955-182
                         29.2%; Score 26; DB 4; Length 8;
  Query Match
  Best Local Similarity 62.5%; Pred. No. 3e+05;
 Matches 5; Conservative 1; Mismatches 2; Indels 0; Gaps
                                                                            0;
           9 QVMMEPQG 16
Qу
             | | | | | | | | |
           1 QQVMTPQG 8
Db
RESULT 25
US-08-475-955-185
; Sequence 185, Application US/08475955
; Patent No. 6641813
; GENERAL INFORMATION:
    APPLICANT: Harley, John
    TITLE OF INVENTION: METHODS AND REAGENTS FOR DIAGNOSIS OF
    TITLE OF INVENTION: AUTOANTIBODIES
    NUMBER OF SEQUENCES: 218
    CORRESPONDENCE ADDRESS:
     ADDRESSEE: Patrea L. Pabst
      STREET: 2800 One Atlantic Center, 1250 West Peachtree Street
      CITY: Atlanta
;
      STATE: GA
;
      COUNTRY: USA
      ZIP: 30309-3450
    COMPUTER READABLE FORM:
      MEDIUM TYPE: Floppy disk
      COMPUTER: IBM PC compatible
      OPERATING SYSTEM: PC-DOS/MS-DOS
      SOFTWARE: PatentIn Release #1.0, Version #1.25
    CURRENT APPLICATION DATA:
      APPLICATION NUMBER: US/08/475,955
      FILING DATE: June 7, 1995
      CLASSIFICATION: 424
    PRIOR APPLICATION DATA:
      APPLICATION NUMBER: 07/867,819
       FILING DATE: April 13, 1992
       CLASSIFICATION: 424
    PRIOR APPLICATION DATA:
      APPLICATION NUMBER: 07/648,205
       FILING DATE: January 31, 1991
     PRIOR APPLICATION DATA:
       APPLICATION NUMBER: 07/472,947
       FILING DATE: January 31, 1990
    ATTORNEY/AGENT INFORMATION:
      NAME: Pabst, Patrea L.
      REGISTRATION NUMBER: 31,284
      REFERENCE/DOCKET NUMBER: OMRF114CIP(2)DIV
    TELECOMMUNICATION INFORMATION:
      TELEPHONE: (404)-873-8794
      TELEFAX: (404)-873-8795
  INFORMATION FOR SEQ ID NO: 185:
     SEQUENCE CHARACTERISTICS:
```

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LENGTH: 8 amino acids
      TYPE: amino acid
      STRANDEDNESS: single
      TOPOLOGY: linear
    MOLECULE TYPE: peptide
US-08-475-955-185
  Query Match
                        29.2%; Score 26; DB 4; Length 8;
  Best Local Similarity 66.7%; Pred. No. 3e+05;
           4; Conservative 1; Mismatches 1; Indels 0; Gaps
 Matches
                                                                          0;
          12 MEPOGD 17
Qy
             1 |||:
Db
           1 MTPQGB 6
RESULT 26
US-08-159-339A-1062
; Sequence 1062, Application US/08159339A
; Patent No. 6037135
; GENERAL INFORMATION:
    APPLICANT: Kubo, Ralph T.
    APPLICANT: Grey, Howard M.
    APPLICANT: Sette, Alessandro
    APPLICANT: Celis, Esteban
    TITLE OF INVENTION: HLA Binding peptides and Their
    TITLE OF INVENTION: Uses
    NUMBER OF SEQUENCES: 1254
    CORRESPONDENCE ADDRESS:
     ADDRESSEE: Townsend and Townsend and Crew LLP
     STREET: Two Embarcadero Center, Eighth Floor
     CITY: San Francisco
      STATE: CA
      COUNTRY: USA
      ZIP: 94111-3834
    COMPUTER READABLE FORM:
     MEDIUM TYPE: Diskette
      COMPUTER: IBM Compatible
      OPERATING SYSTEM: DOS
      SOFTWARE: FastSEQ for Windows Version 2.0
    CURRENT APPLICATION DATA:
      APPLICATION NUMBER: US/08/159,339A
      FILING DATE: 29-NOV-1993
      CLASSIFICATION: 424
    PRIOR APPLICATION DATA:
     APPLICATION NUMBER: US 07/926,666
      FILING DATE: 07-AUG-1992
      APPLICATION NUMBER: US 08/027,746
      FILING DATE: 05-MAR-1993
      APPLICATION NUMBER: US 08/103,396
      FILING DATE: 06-AUG-1993
    ATTORNEY/AGENT INFORMATION:
     NAME: Weber, Ellen Lauver
      REGISTRATION NUMBER: 32,762
      REFERENCE/DOCKET NUMBER: 018623-005030US
    TELECOMMUNICATION INFORMATION:
      TELEPHONE: (415) 576-0200
```

```
TELEFAX: (415) 576-0300
      TELEX:
  INFORMATION FOR SEQ ID NO: 1062:
    SEQUENCE CHARACTERISTICS:
      LENGTH: 9 amino acids
      TYPE: amino acid
       STRANDEDNESS: single
      TOPOLOGY: linear
    MOLECULE TYPE: peptide
US-08-159-339A-1062
 Query Match 29.2%; Score 26; DB 3; Length 9; Best Local Similarity 80.0%; Pred. No. 3e+05;
 Matches
          4; Conservative 1; Mismatches
                                                 0; Indels 0; Gaps
                                                                               0;
           13 EPQGD 17
Qу
              : | | | |
            1 QPQGD 5
Db
RESULT 27
US-08-737-109-5
; Sequence 5, Application US/08737109
; Patent No. 6455688
  GENERAL INFORMATION:
    APPLICANT: SLABAS, Antoni Ryszard APPLICANT: ELBOROUGH, Kieran Michael
    APPLICANT: BRIGHT, Simon William Jonathan
     APPLICANT: FENTEM, Philip Anthony
     TITLE OF INVENTION: Plant Gene Specifying Acetyl Coenzyme A
    TITLE OF INVENTION: Carboxylase and Transformed Plants Containing Same
    NUMBER OF SEQUENCES: 32
    CORRESPONDENCE ADDRESS:
       ADDRESSEE: Pillsbury Madison & Sutro, L.L.P.
       STREET: 1100 New York Avenue, N.W.
      CITY: Washington
      STATE: D.C.
       COUNTRY: U.S.A.
    COMPUTER READABLE FORM:
      MEDIUM TYPE: 3.5 inch diskette
       COMPUTER: IBM PC compatible
       OPERATING SYSTEM: PC-DOS/MS-DOS
       SOFTWARE: MS Word
     CURRENT APPLICATION DATA:
     APPLICATION NUMBER: US/08/737,109
      FILING DATE: 21-OCT-1996
      CLASSIFICATION: 800
     PRIOR APPLICATION DATA:
       APPLICATION NUMBER: PCT/GB94/00846
       FILING DATE: 02-MAY-1994
   INFORMATION FOR SEQ ID NO: 5:
     SEQUENCE CHARACTERISTICS:
       LENGTH: 9 amino acids
       TYPE: amino acid
       TOPOLOGY: linear
     MOLECULE TYPE: peptide
    HYPOTHETICAL: NO
```

```
FRAGMENT TYPE: internal
    ORIGINAL SOURCE:
;
      ORGANISM: Avena sativa
US-08-737-109-5
                         29.2%; Score 26; DB 4; Length 9;
 Query Match
 Best Local Similarity 66.7%; Pred. No. 3e+05;
 Matches 4; Conservative 2; Mismatches 0; Indels 0; Gaps
                                                                           0;
         11 MMEPQG 16
Qy
             ::||||
           2 VLEPQG 7
RESULT 28
5204326-100
; Patent No. 5204326
    APPLICANT: FUJII, SETSURO; YAMAMOTO, YOSHIHITO; SHIMIZU, FUMIO
; INAI, MASATOSHI; KINOSHITA, NAOSUMI; NAKAMURA, SHIZUO; HIROHASHI,
;MITSURU; SAKAMOTO, TAKASHI;TSUTSUMI, KAZUHIKO;SHIRASAKA, TETSUHIKO
    TITLE OF INVENTION: POLYPEPTIDE DERIVATIVES AND CALCIUM
;METABOLISM IMPROVING AGENT
    NUMBER OF SEQUENCES: 147
    CURRENT APPLICATION DATA:
      APPLICATION NUMBER: US/07/493,359
      FILING DATE: 14-MAR-1990
;SEO ID NO:100:
      LENGTH: 9
5204326-100
                         29.2%; Score 26; DB 6; Length 9;
  Query Match
  Best Local Similarity 100.0%; Pred. No. 3e+05;
           5; Conservative 0; Mismatches
                                                0; Indels 0; Gaps
                                                                           0;
 Matches
           1 LQTPQ 5
Qу
             11111
Db
           3 LQTPQ 7
RESULT 29
US-08-159-339A-971
; Sequence 971, Application US/08159339A
; Patent No. 6037135
; GENERAL INFORMATION:
    APPLICANT: Kubo, Ralph T.
    APPLICANT: Grey, Howard M.
    APPLICANT: Sette, Alessandro
    APPLICANT: Celis, Esteban
    TITLE OF INVENTION: HLA Binding peptides and Their
    TITLE OF INVENTION: Uses
    NUMBER OF SEQUENCES: 1254
    CORRESPONDENCE ADDRESS:
     ADDRESSEE: Townsend and Townsend and Crew LLP
      STREET: Two Embarcadero Center, Eighth Floor
      CITY: San Francisco
      STATE: CA
      COUNTRY: USA
```

```
ZIP: 94111-3834
    COMPUTER READABLE FORM:
      MEDIUM TYPE: Diskette
      COMPUTER: IBM Compatible
      OPERATING SYSTEM: DOS
      SOFTWARE: FastSEQ for Windows Version 2.0
    CURRENT APPLICATION DATA:
     APPLICATION NUMBER: US/08/159,339A
     FILING DATE: 29-NOV-1993
     CLASSIFICATION: 424
    PRIOR APPLICATION DATA:
    APPLICATION NUMBER: US 07/926,666
      FILING DATE: 07-AUG-1992
     APPLICATION NUMBER: US 08/027,746
     FILING DATE: 05-MAR-1993
     APPLICATION NUMBER: US 08/103,396
      FILING DATE: 06-AUG-1993
    ATTORNEY/AGENT INFORMATION:
     NAME: Weber, Ellen Lauver
      REGISTRATION NUMBER: 32,762
      REFERENCE/DOCKET NUMBER: 018623-005030US
    TELECOMMUNICATION INFORMATION:
      TELEPHONE: (415) 576-0200
      TELEFAX: (415) 576-0300
      TELEX:
  INFORMATION FOR SEQ ID NO: 971:
    SEQUENCE CHARACTERISTICS:
      LENGTH: 10 amino acids
      TYPE: amino acid
      STRANDEDNESS: single
      TOPOLOGY: linear
    MOLECULE TYPE: peptide
US-08-159-339A-971
 Query Match
                         29.2%; Score 26; DB 3; Length 10;
 Best Local Similarity 80.0%; Pred. No. 3.3e+02;
 Matches 4; Conservative 1; Mismatches 0; Indels 0; Gaps
                                                                           0;
          13 EPQGD 17
Qy
             : | | | |
Dh
           5 QPQGD 9
RESULT 30
US-08-159-339A-1070
; Sequence 1070, Application US/08159339A
; Patent No. 6037135
 GENERAL INFORMATION:
    APPLICANT: Kubo, Ralph T.
    APPLICANT: Grey, Howard M.
APPLICANT: Sette, Alessandro
    APPLICANT: Celis, Esteban
    TITLE OF INVENTION: HLA Binding peptides and Their
    TITLE OF INVENTION: Uses
   NUMBER OF SEQUENCES: 1254
    CORRESPONDENCE ADDRESS:
      ADDRESSEE: Townsend and Townsend and Crew LLP
```

```
STREET: Two Embarcadero Center, Eighth Floor
      CITY: San Francisco
      STATE: CA
      COUNTRY: USA
      ZIP: 94111-3834
    COMPUTER READABLE FORM:
      MEDIUM TYPE: Diskette
      COMPUTER: IBM Compatible
      OPERATING SYSTEM: DOS
      SOFTWARE: FastSEQ for Windows Version 2.0
    CURRENT APPLICATION DATA:
      APPLICATION NUMBER: US/08/159,339A
      FILING DATE: 29-NOV-1993
      CLASSIFICATION: 424
    PRIOR APPLICATION DATA:
      APPLICATION NUMBER: US 07/926,666
      FILING DATE: 07-AUG-1992
      APPLICATION NUMBER: US 08/027,746
      FILING DATE: 05-MAR-1993
      APPLICATION NUMBER: US 08/103,396
      FILING DATE: 06-AUG-1993
    ATTORNEY/AGENT INFORMATION:
      NAME: Weber, Ellen Lauver
      REGISTRATION NUMBER: 32,762
      REFERENCE/DOCKET NUMBER: 018623-005030US
    TELECOMMUNICATION INFORMATION:
      TELEPHONE: (415) 576-0200
      TELEFAX: (415) 576-0300
      TELEX:
  INFORMATION FOR SEQ ID NO: 1070:
    SEQUENCE CHARACTERISTICS:
      LENGTH: 10 amino acids
      TYPE: amino acid
      STRANDEDNESS: single
      TOPOLOGY: linear
    MOLECULE TYPE: peptide
US-08-159-339A-1070
 Query Match
                         29.2%; Score 26; DB 3; Length 10;
 Best Local Similarity 80.0%; Pred. No. 3.3e+02;
 Matches 4; Conservative 1; Mismatches 0; Indels 0; Gaps
                                                                           0;
          13 EPQGD 17
Qу
             : | | | |
           1 QPQGD 5
Db
RESULT 31
US-09-023-173-9
; Sequence 9, Application US/09023173
; Patent No. 6066781
; GENERAL INFORMATION:
    APPLICANT: Sutliff, Thomas D.
    APPLICANT: Rodriguez, Raymond L.
    TITLE OF INVENTION: Production of Mature Proteins
    TITLE OF INVENTION: in Plants
    NUMBER OF SEQUENCES: 23
```

```
CORRESPONDENCE ADDRESS:
      ADDRESSEE: Dehlinger & Associates
      STREET: 350 Cambridge Ave., Suite 250
      CITY: Palo Alto
      STATE: CA
      COUNTRY: USA
;
      ZIP: 94306
    COMPUTER READABLE FORM:
      MEDIUM TYPE: Diskette
;
      COMPUTER: IBM Compatible
      OPERATING SYSTEM: DOS
      SOFTWARE: FastSEQ for Windows Version 2.0
;
    CURRENT APPLICATION DATA:
;
     APPLICATION NUMBER: US/09/023,173
      FILING DATE: 13-FEB-1998
      CLASSIFICATION: 435
    PRIOR APPLICATION DATA:
      APPLICATION NUMBER: 60/038,168
      FILING DATE: 13-FEB-1997
;
    ATTORNEY/AGENT INFORMATION:
;
     NAME: Petithory, Joanne R
      REGISTRATION NUMBER: P42995
      REFERENCE/DOCKET NUMBER: 0665-0007.30
    TELECOMMUNICATION INFORMATION:
      TELEPHONE: 650-324-0880
      TELEFAX: 650-324-0960
;
  INFORMATION FOR SEO ID NO: 9:
    SEQUENCE CHARACTERISTICS:
;
     LENGTH: 13 amino acids
      TYPE: amino acid
      STRANDEDNESS: single
      TOPOLOGY: linear
    MOLECULE TYPE: peptide
    IMMEDIATE SOURCE:
      CLONE: N-terminal amino acid sequence of mature AAT
US-09-023-173-9
  Query Match
                         29.2%; Score 26; DB 3; Length 13;
  Best Local Similarity 80.0%; Pred. No. 4.5e+02;
 Matches
            4; Conservative 1; Mismatches 0; Indels 0; Gaps
                                                                           0;
          13 EPQGD 17
Qу
             :1111
           2 DPQGD 6
Db
RESULT 32
US-09-023-339-7
; Sequence 7, Application US/09023339
; Patent No. 6127145
; GENERAL INFORMATION:
    APPLICANT: Sutliff, Thomas D.
    APPLICANT: Rodriguez, Raymond L.
    TITLE OF INVENTION: Production of `1-Antitrypsin
    TITLE OF INVENTION: in Plants
    NUMBER OF SEQUENCES: 22
    CORRESPONDENCE ADDRESS:
```

```
ADDRESSEE: Dehlinger & Associates
       STREET: P.O. Box 60850
       CITY: Palo Alto
       STATE: CA
       COUNTRY: USA
       ZIP: 94306
;
    COMPUTER READABLE FORM:
      MEDIUM TYPE: Diskette
       COMPUTER: IBM Compatible
       OPERATING SYSTEM: DOS
       SOFTWARE: FastSEQ for Windows Version 2.0
     CURRENT APPLICATION DATA:
;
       APPLICATION NUMBER: US/09/023,339
;
       FILING DATE: 13-FEB-1998
     PRIOR APPLICATION DATA:
       APPLICATION NUMBER: 60/037,991
       FILING DATE: 13-FEB-1997
    ATTORNEY/AGENT INFORMATION:
      NAME: Petithory, Joanne R
       REGISTRATION NUMBER: P42,995
       REFERENCE/DOCKET NUMBER: 0665-0003.30
     TELECOMMUNICATION INFORMATION:
       TELEPHONE: 650-324-0880
       TELEFAX: 650-324-0960
   INFORMATION FOR SEQ ID NO:
     SEQUENCE CHARACTERISTICS:
       LENGTH: 13 amino acids
;
       TYPE: amino acid
;
       TOPOLOGY: linear
    MOLECULE TYPE: peptide
     IMMEDIATE SOURCE:
       CLONE: N-terminal sequence of plant-produced mature AAT
US-09-023-339-7
  Query Match
                         29.2%; Score 26; DB 3; Length 13;
                         80.0%; Pred. No. 4.5e+02;
  Best Local Similarity
  Matches
           4; Conservative
                               1; Mismatches 0; Indels 0; Gaps
                                                                            0;
           13 EPQGD 17
Qу
              : | | | |
            2 DPQGD 6
Db
RESULT 33
US-08-855-744-2
; Sequence 2, Application US/08855744
; Patent No. 6685930
  GENERAL INFORMATION:
     APPLICANT: Chang, Tse Wen
     TITLE OF INVENTION: METHODS AND SUBSTANCES FOR RECRUITING
     TITLE OF INVENTION: THERAPEUTIC AGENTS TO SOLID TISSUES
;
    NUMBER OF SEQUENCES: 3
     CORRESPONDENCE ADDRESS:
       ADDRESSEE: Tanox Biosystems, Inc.
       STREET: 10301 Stella Link Rd.
      CITY: Houston
      STATE: Texas
```

```
COUNTRY: USA
      ZIP: 77025
    COMPUTER READABLE FORM:
      MEDIUM TYPE: 3.5" Hi Density Diskette
      COMPUTER: IBM PS/2
;
      OPERATING SYSTEM: DOS, Version 3.30
;
      SOFTWARE: Wordperfect 5.1
    CURRENT APPLICATION DATA:
    APPLICATION NUMBER: US/08/855,744
     FILING DATE: 08-May-1997
     CLASSIFICATION: 424
    PRIOR APPLICATION DATA:
;
    APPLICATION NUMBER: US/08/264,649
;
      FILING DATE:
     APPLICATION NUMBER: US/07/675,654
      FILING DATE:
    ATTORNEY/AGENT INFORMATION:
     NAME: Mirabel, Eric P.
      REGISTRATION NUMBER: 31,211
;
      REFERENCE/DOCKET NUMBER: TNX91-2
    TELECOMMUNICATION INFORMATION:
      TELEPHONE: 713-664-2288
      TELEFAX: 713-664-8914
  INFORMATION FOR SEQ ID NO: 2:
    SEQUENCE CHARACTERISTICS:
      LENGTH: 14 amino acids
      TYPE: amino acid
      TOPOLOGY: Linear
US-08-855-744-2
                        29.2%; Score 26; DB 4; Length 14;
  Query Match
  Best Local Similarity 62.5%; Pred. No. 4.8e+02;
           5; Conservative 1; Mismatches 2; Indels 0; Gaps
 Matches
                                                                           0;
           3 TPQPLLQV 10
Qу
            -11 + 1 = 11
Db
           4 TPSPGIQV 11
RESULT 34
5204326-36
; Patent No. 5204326
    APPLICANT: FUJII, SETSURO; YAMAMOTO, YOSHIHITO; SHIMIZU, FUMIO
; INAI, MASATOSHI; KINOSHITA, NAOSUMI; NAKAMURA, SHIZUO; HIROHASHI,
;MITSURU; SAKAMOTO, TAKASHI;TSUTSUMI, KAZUHIKO;SHIRASAKA, TETSUHIKO
    TITLE OF INVENTION: POLYPEPTIDE DERIVATIVES AND CALCIUM
;METABOLISM IMPROVING AGENT
    NUMBER OF SEQUENCES: 147
    CURRENT APPLICATION DATA:
      APPLICATION NUMBER: US/07/493,359
      FILING DATE: 14-MAR-1990
; SEQ ID NO: 36:
      LENGTH: 14
5204326-36
  Query Match
                         29.2%; Score 26; DB 6; Length 14;
  Best Local Similarity 100.0%; Pred. No. 4.8e+02;
```

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Matches 5; Conservative 0; Mismatches 0; Indels 0; Gaps
                                                                          0;
Qу
           1 LQTPQ 5
             Db
           2 LQTPQ 6
RESULT 35
5204326-95
; Patent No. 5204326
    APPLICANT: FUJII, SETSURO; YAMAMOTO, YOSHIHITO; SHIMIZU, FUMIO
; INAI, MASATOSHI; KINOSHITA, NAOSUMI; NAKAMURA, SHIZUO; HIROHASHI,
;MITSURU; SAKAMOTO, TAKASHI;TSUTSUMI, KAZUHIKO;SHIRASAKA, TETSUHIKO
    TITLE OF INVENTION: POLYPEPTIDE DERIVATIVES AND CALCIUM
;METABOLISM IMPROVING AGENT
    NUMBER OF SEQUENCES: 147
    CURRENT APPLICATION DATA:
      APPLICATION NUMBER: US/07/493,359
      FILING DATE: 14-MAR-1990
;SEQ ID NO:95:
      LENGTH: 14
5204326-95
 Query Match
                         29.2%; Score 26; DB 6; Length 14;
  Best Local Similarity 100.0%; Pred. No. 4.8e+02;
 Matches 5; Conservative 0; Mismatches 0; Indels 0; Gaps
                                                                          0;
           1 LQTPQ 5
Qу
            2 LQTPQ 6
Db
RESULT 36
US-08-096-044C-14
; Sequence 14, Application US/08096044C
; Patent No. 6153192
; GENERAL INFORMATION:
    APPLICANT: Kopetzki, Erhard, Klein, Christian
    TITLE OF INVENTION: PEPTIDES WITH CHARACTERISTIC
    TITLE OF INVENTION: ANTIGENIC DETERMINANTS FOR
    TITLE OF INVENTION: 1-MICROGLOBULIN
    NUMBER OF SEQUENCES: 18
    CORRESPONDENCE ADDRESS:
     ADDRESSEE: Felfe & Lynch
      STREET: 805 Third Avenue
      CITY: New York
      STATE: New York
      COUNTRY: USA
      ZIP: 10022
    COMPUTER READABLE FORM:
      MEDIUM TYPE: 3.5 inch 1.44 Mb storage diskette
      COMPUTER: IBM PS/2
      OPERATING SYSTEM: PC-DOS
      SOFTWARE: Wordperfect
    CURRENT APPLICATION DATA:
      APPLICATION NUMBER: US/08/096,044C
      FILING DATE: July 22, 1993
```

```
CLASSIFICATION: 424
    PRIOR APPLICATION DATA:
      APPLICATION NUMBER: DE 40 24 919
;
      FILING DATE: August 6, 1990
;
    ATTORNEY/AGENT INFORMATION:
;
      NAME: Bauer, John A.
      REGISTRATION NUMBER: 32,554
      REFERENCE/DOCKET NUMBER: HUBR 1000.1-PFF/JAB
    TELECOMMUNICATION INFORMATION:
      TELEPHONE: (212) 688-9200
      TELEFAX: (212) 838-3884
  INFORMATION FOR SEQ ID NO: 14:
    SEQUENCE CHARACTERISTICS:
      LENGTH: 15
      TYPE: amino acid
      STRANDEDNESS: single
      TOPOLOGY: linear
US-08-096-044C-14
                         29.2%; Score 26; DB 3; Length 15;
  Query Match
  Best Local Similarity 57.1%; Pred. No. 5.2e+02;
           4; Conservative 2; Mismatches 1; Indels 0; Gaps
 Matches
                                                                           0;
           2 QTPQPLL 8
Qу
             1 1:1:1
           6 QEPEPIL 12
Db
RESULT 37
5204326-37
; Patent No. 5204326
    APPLICANT: FUJII, SETSURO; YAMAMOTO, YOSHIHITO; SHIMIZU, FUMIO
; INAI, MASATOSHI; KINOSHITA, NAOSUMI; NAKAMURA, SHIZUO; HIROHASHI,
;MITSURU; SAKAMOTO, TAKASHI;TSUTSUMI, KAZUHIKO;SHIRASAKA, TETSUHIKO
    TITLE OF INVENTION: POLYPEPTIDE DERIVATIVES AND CALCIUM
;METABOLISM IMPROVING AGENT
    NUMBER OF SEQUENCES: 147
    CURRENT APPLICATION DATA:
      APPLICATION NUMBER: US/07/493,359
      FILING DATE: 14-MAR-1990
;SEQ ID NO:37:
      LENGTH: 15
5204326-37
  Query Match
                         29.2%; Score 26; DB 6; Length 15;
  Best Local Similarity 100.0%; Pred. No. 5.2e+02;
           5; Conservative 0; Mismatches
                                                0; Indels
                                                               0; Gaps
                                                                           0;
           1 LOTPO 5
Qу
             3 LOTPO 7
Db
RESULT 38
5204326-40
; Patent No. 5204326
    APPLICANT: FUJII, SETSURO; YAMAMOTO, YOSHIHITO; SHIMIZU, FUMIO
```

```
; INAI, MASATOSHI; KINOSHITA, NAOSUMI; NAKAMURA, SHIZUO; HIROHASHI,
;MITSURU; SAKAMOTO, TAKASHI;TSUTSUMI, KAZUHIKO;SHIRASAKA, TETSUHIKO
    TITLE OF INVENTION: POLYPEPTIDE DERIVATIVES AND CALCIUM
; METABOLISM IMPROVING AGENT
    NUMBER OF SEQUENCES: 147
    CURRENT APPLICATION DATA:
      APPLICATION NUMBER: US/07/493,359
       FILING DATE: 14-MAR-1990
;SEQ ID NO:40:
      LENGTH: 15
5204326-40
  Query Match
                         29.2%; Score 26; DB 6; Length 15;
  Best Local Similarity 100.0%; Pred. No. 5.2e+02;
            5; Conservative
                                0; Mismatches
                                                0; Indels 0; Gaps
                                                                             0;
           1 LQTPQ 5
Qу
             +11111
           3 LQTPQ 7
Db
RESULT 39
5204326-71
; Patent No. 5204326
    APPLICANT: FUJII, SETSURO; YAMAMOTO, YOSHIHITO; SHIMIZU, FUMIO
; INAI, MASATOSHI; KINOSHITA, NAOSUMI; NAKAMURA, SHIZUO; HIROHASHI,
;MITSURU; SAKAMOTO, TAKASHI;TSUTSUMI, KAZUHIKO;SHIRASAKA, TETSUHIKO
     TITLE OF INVENTION: POLYPEPTIDE DERIVATIVES AND CALCIUM
;METABOLISM IMPROVING AGENT
    NUMBER OF SEQUENCES: 147
     CURRENT APPLICATION DATA:
       APPLICATION NUMBER: US/07/493,359
      FILING DATE: 14-MAR-1990
;SEQ ID NO:71:
      LENGTH: 15
5204326-71
  Query Match
                         29.2%; Score 26; DB 6; Length 15;
  Best Local Similarity 100.0%; Pred. No. 5.2e+02;
           5; Conservative 0; Mismatches 0; Indels
                                                                 0; Gaps
                                                                             0;
  Matches
            1 LQTPQ 5
Qу
             3 LQTPQ 7
Db
RESULT 40
5204326-74
; Patent No. 5204326
     APPLICANT: FUJII, SETSURO; YAMAMOTO, YOSHIHITO; SHIMIZU, FUMIO
; INAI, MASATOSHI; KINOSHITA, NAOSUMI; NAKAMURA, SHIZUO; HIROHASHI,
;MITSURU; SAKAMOTO, TAKASHI;TSUTSUMI, KAZUHIKO;SHIRASAKA, TETSUHIKO
     TITLE OF INVENTION: POLYPEPTIDE DERIVATIVES AND CALCIUM
; METABOLISM IMPROVING AGENT
     NUMBER OF SEQUENCES: 147
     CURRENT APPLICATION DATA:
       APPLICATION NUMBER: US/07/493,359
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FILING DATE: 14-MAR-1990
;SEQ ID NO:74:
      LENGTH: 15
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Qу
             3 LOTPO 7
Db
RESULT 41
US-07-700-526-8
; Sequence 8, Application US/07700526
; Patent No. 5166133
  GENERAL INFORMATION:
    APPLICANT: Houston, L. L.
    APPLICANT: Liu, David Y.
    APPLICANT: Kaymakcalan, Zehra
    TITLE OF INVENTION: Method for Inhibiting Adhesion of White
    TITLE OF INVENTION: Blood Cells to Endothelial Cells
    NUMBER OF SEQUENCES: 18
    CORRESPONDENCE ADDRESS:
;
      ADDRESSEE: Cetus Corporation
;
      STREET: 1400 Fifty-Third Street
;
     CITY: Emeryville
     STATE: CA
     COUNTRY: USA
     ZIP: 94608
    COMPUTER READABLE FORM:
      MEDIUM TYPE: Floppy disk
      COMPUTER: IBM PC compatible
      OPERATING SYSTEM: PC-DOS/MS-DOS
      SOFTWARE: PatentIn Release #1.0, Version #1.25
    CURRENT APPLICATION DATA:
     APPLICATION NUMBER: US/07/700,526
      FILING DATE: 19910816
      CLASSIFICATION: 424
    ATTORNEY/AGENT INFORMATION:
      NAME: McGarrigle Jr., Philip L.
      REGISTRATION NUMBER: 31,395
      REFERENCE/DOCKET NUMBER: 2600.1
    TELECOMMUNICATION INFORMATION:
      TELEPHONE: (415) 420-3217
      TELEFAX: (415) 658-5239
      TELEX: 4992659
  INFORMATION FOR SEQ ID NO: 8:
    SEQUENCE CHARACTERISTICS:
      LENGTH: 16 amino acids
      TYPE: AMINO ACID
      STRANDEDNESS: single
      TOPOLOGY: linear
    MOLECULE TYPE: protein
US-07-700-526-8
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  Best Local Similarity 57.1%; Pred. No. 5.6e+02;
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           4; Conservative 2; Mismatches 1; Indels 0; Gaps
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Qy
             1 1:1:1
           8 OEPEPIL 14
Db
RESULT 42
US-08-096-044C-13
; Sequence 13, Application US/08096044C
; Patent No. 6153192
  GENERAL INFORMATION:
    APPLICANT: Kopetzki, Erhard, Klein, Christian
    TITLE OF INVENTION: PEPTIDES WITH CHARACTERISTIC
    TITLE OF INVENTION: ANTIGENIC DETERMINANTS FOR
    TITLE OF INVENTION: 1-MICROGLOBULIN
    NUMBER OF SEQUENCES: 18
    CORRESPONDENCE ADDRESS:
     ADDRESSEE: Felfe & Lynch
      STREET: 805 Third Avenue
      CITY: New York
      STATE: New York
      COUNTRY: USA
;
      ZIP: 10022
;
    COMPUTER READABLE FORM:
      MEDIUM TYPE: 3.5 inch 1.44 Mb storage diskette
      COMPUTER: IBM PS/2
      OPERATING SYSTEM: PC-DOS
      SOFTWARE: Wordperfect
    CURRENT APPLICATION DATA:
      APPLICATION NUMBER: US/08/096,044C
      FILING DATE: July 22, 1993
      CLASSIFICATION: 424
    PRIOR APPLICATION DATA:
      APPLICATION NUMBER: DE 40 24 919
      FILING DATE: August 6, 1990
    ATTORNEY/AGENT INFORMATION:
      NAME: Bauer, John A.
      REGISTRATION NUMBER: 32,554
      REFERENCE/DOCKET NUMBER: HUBR 1000.1-PFF/JAB
    TELECOMMUNICATION INFORMATION:
      TELEPHONE: (212) 688-9200
      TELEFAX: (212) 838-3884
  INFORMATION FOR SEQ ID NO: 13:
    SEQUENCE CHARACTERISTICS:
      LENGTH: 16
      TYPE: amino acid
      STRANDEDNESS: single
      TOPOLOGY: linear
US-08-096-044C-13
  Query Match
                         29.2%; Score 26; DB 3; Length 16;
 Best Local Similarity 57.1%; Pred. No. 5.6e+02;
  Matches 4; Conservative 2; Mismatches 1; Indels 0; Gaps
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             1 1:1:1
           7 QEPEPIL 13
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US-08-096-044C-15
; Sequence 15, Application US/08096044C
; Patent No. 6153192
  GENERAL INFORMATION:
    APPLICANT: Kopetzki, Erhard, Klein, Christian
    TITLE OF INVENTION: PEPTIDES WITH CHARACTERISTIC
    TITLE OF INVENTION: ANTIGENIC DETERMINANTS FOR
    TITLE OF INVENTION: 1-MICROGLOBULIN
    NUMBER OF SEQUENCES: 18
    CORRESPONDENCE ADDRESS:
      ADDRESSEE: Felfe & Lynch
      STREET: 805 Third Avenue
      CITY: New York
      STATE: New York
      COUNTRY: USA
      ZIP: 10022
    COMPUTER READABLE FORM:
      MEDIUM TYPE: 3.5 inch 1.44 Mb storage diskette
      COMPUTER: IBM PS/2
      OPERATING SYSTEM: PC-DOS
      SOFTWARE: Wordperfect
    CURRENT APPLICATION DATA:
      APPLICATION NUMBER: US/08/096,044C
      FILING DATE: July 22, 1993
      CLASSIFICATION: 424
    PRIOR APPLICATION DATA:
      APPLICATION NUMBER: DE 40 24 919
      FILING DATE: August 6, 1990
    ATTORNEY/AGENT INFORMATION:
      NAME: Bauer, John A.
       REGISTRATION NUMBER: 32,554
       REFERENCE/DOCKET NUMBER: HUBR 1000.1-PFF/JAB
     TELECOMMUNICATION INFORMATION:
      TELEPHONE: (212) 688-9200
       TELEFAX: (212) 838-3884
  INFORMATION FOR SEQ ID NO: 15:
    SEQUENCE CHARACTERISTICS:
      LENGTH: 16
       TYPE: amino acid
       STRANDEDNESS: single
       TOPOLOGY: linear
US-08-096-044C-15
                         29.2%; Score 26; DB 3; Length 16;
  Query Match
  Best Local Similarity 57.1%; Pred. No. 5.6e+02;
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          4; Conservative 2; Mismatches 1; Indels
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           2 QTPQPLL 8
Qу
             1 1:1:1
Db
           7 QEPEPIL 13
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RESULT 44
PCT-US92-03132-8
; Sequence 8, Application PC/TUS9203132
  GENERAL INFORMATION:
    APPLICANT: Houston, L. L.
    APPLICANT: Liu, David Y.
    APPLICANT: Kaymakcalan, Zehra
    TITLE OF INVENTION: Method for Inhibiting Adhesion of White
    TITLE OF INVENTION: Blood Cells to Endothelial Cells
    NUMBER OF SEQUENCES: 18
;
    CORRESPONDENCE ADDRESS:
      ADDRESSEE: Cetus Corporation
      STREET: 1400 Fifty-Third Street
;
      CITY: Emeryville
      STATE: CA
      COUNTRY: USA
;
      ZIP: 94608
;
    COMPUTER READABLE FORM:
      MEDIUM TYPE: Floppy disk
      COMPUTER: IBM PC compatible
      OPERATING SYSTEM: PC-DOS/MS-DOS
;
      SOFTWARE: PatentIn Release #1.0, Version #1.25
    CURRENT APPLICATION DATA:
;
      APPLICATION NUMBER: PCT/US92/03132
;
      FILING DATE: 19920416
;
      CLASSIFICATION:
    PRIOR APPLICATION DATA:
      APPLICATION NUMBER: US/07/700,526
      FILING DATE: 14-MAY-1991
    ATTORNEY/AGENT INFORMATION:
      NAME: McGarrigle Jr., Philip L.
      REGISTRATION NUMBER: 31,395
      REFERENCE/DOCKET NUMBER:
    TELECOMMUNICATION INFORMATION:
      TELEPHONE: (415) 420-3217
      TELEFAX: (415) 658-5239
       TELEX: 4992659
   INFORMATION FOR SEQ ID NO: 8:
    SEQUENCE CHARACTERISTICS:
       LENGTH: 16 amino acids
      TYPE: AMINO ACID
      STRANDEDNESS: single
      TOPOLOGY: linear
    MOLECULE TYPE: protein
PCT-US92-03132-8
  Query Match
                         29.2%; Score 26; DB 5; Length 16;
                         57.1%; Pred. No. 5.6e+02;
  Best Local Similarity
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Qy
              1 1:1:1
Db
            8 QEPEPIL 14
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RESULT 45
5204326-96
; Patent No. 5204326
    APPLICANT: FUJII, SETSURO; YAMAMOTO, YOSHIHITO; SHIMIZU, FUMIO
; INAI, MASATOSHI; KINOSHITA, NAOSUMI; NAKAMURA, SHIZUO; HIROHASHI,
;MITSURU; SAKAMOTO, TAKASHI;TSUTSUMI, KAZUHIKO;SHIRASAKA, TETSUHIKO
    TITLE OF INVENTION: POLYPEPTIDE DERIVATIVES AND CALCIUM
;METABOLISM IMPROVING AGENT
    NUMBER OF SEQUENCES: 147
    CURRENT APPLICATION DATA:
      APPLICATION NUMBER: US/07/493,359
      FILING DATE: 14-MAR-1990
;SEQ ID NO:96:
      LENGTH: 16
5204326-96
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                         29.2%; Score 26; DB 6; Length 16;
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Qу
             Db
           4 LQTPQ 8
RESULT 46
US-08-675-921B-7
; Sequence 7, Application US/08675921B
; Patent No. 5863728
  GENERAL INFORMATION:
    APPLICANT: John Siu-Cheong Ho, John T. Loh, Melvin
    APPLICANT: Schindler and John L. Wang
    TITLE OF INVENTION: DNA Encoding Carbohydrate
    TITLE OF INVENTION: Binding Protein and Biological
    TITLE OF INVENTION: Materials Derived Therefrom
    NUMBER OF SEQUENCES: 9
    CORRESPONDENCE ADDRESS:
     ADDRESSEE: Ian C. McLeod
      STREET: 2190 Commons Parkway
      CITY: Okemos
      STATE: Michigan
;
      COUNTRY: USA
      ZIP: 48864
    COMPUTER READABLE FORM:
      MEDIUM TYPE: Diskette 5.25 inch, 360
      MEDIUM TYPE: Kb storage
      COMPUTER: IBM compatible
      OPERATING SYSTEM: MS-DOS (version 3.3)
      SOFTWARE: Wordperfect 5.1
    CURRENT APPLICATION DATA:
      APPLICATION NUMBER: US/08/675,921B
      FILING DATE: 06/05/96
      CLASSIFICATION: 435
    PRIOR APPLICATION DATA:
      APPLICATION NUMBER:
       FILING DATE:
    ATTORNEY/AGENT INFORMATION:
```

```
NAME: Ian C. McLeod
      REGISTRATION NUMBER: 20,931
      REFERENCE/DOCKET NUMBER: MSU 4.1-265
    TELECOMMUNICATION INFORMATION:
      TELEPHONE: (517) 347-4100
      TELEFAX: (517) 347-4103
      TELEX: No. 5863728e
  INFORMATION FOR SEQ ID NO: 7:
    SEOUENCE CHARACTERISTICS:
      LENGTH: 17
      TYPE: Amino Acid
      STRANDEDNESS: Single
      TOPOLOGY: Linear
    MOLECULE TYPE:
      DESCRIPTION: Peptide
    HYPOTHETICAL: No
    ANTI-SENSE: No
    FRAGMENT TYPE: N/A
    ORIGINAL SOURCE: N/A
      ORGANISM: N/A
      STRAIN: N/A
      INDIVIDUAL ISOLATE: N/A
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      HAPLOTYPE: N/A
      TISSUE TYPE: N/A
      CELL TYPE: N/A
      CELL LINE: N/A
      ORGANELLE: N/A
    IMMEDIATE SOURCE:
    POSITION IN GENOME: N/A
    FEATURE:
      NAME/KEY: deduced partial amino acid
      NAME/KEY: sequence of BJ38 DNA
      LOCATION: N/A
      IDENTIFICATION METHOD: N/A
      OTHER INFORMATION: N/A
    PUBLICATION INFORMATION: N/A
US-08-675-921B-7
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RESULT 47
5204326-97
; Patent No. 5204326
    APPLICANT: FUJII, SETSURO; YAMAMOTO, YOSHIHITO; SHIMIZU, FUMIO
; INAI, MASATOSHI; KINOSHITA, NAOSUMI; NAKAMURA, SHIZUO; HIROHASHI,
;MITSURU; SAKAMOTO, TAKASHI;TSUTSUMI, KAZUHIKO;SHIRASAKA, TETSUHIKO
    TITLE OF INVENTION: POLYPEPTIDE DERIVATIVES AND CALCIUM
;METABOLISM IMPROVING AGENT
    NUMBER OF SEQUENCES: 147
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CURRENT APPLICATION DATA:
      APPLICATION NUMBER: US/07/493,359
      FILING DATE: 14-MAR-1990
; SEQ ID NO:97:
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5204326-97
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RESULT 48
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; Patent No. 5204326
    APPLICANT: FUJII, SETSURO; YAMAMOTO, YOSHIHITO; SHIMIZU, FUMIO
; INAI, MASATOSHI; KINOSHITA, NAOSUMI; NAKAMURA, SHIZUO; HIROHASHI,
;MITSURU; SAKAMOTO, TAKASHI;TSUTSUMI, KAZUHIKO;SHIRASAKA, TETSUHIKO
    TITLE OF INVENTION: POLYPEPTIDE DERIVATIVES AND CALCIUM
;METABOLISM IMPROVING AGENT
    NUMBER OF SEQUENCES: 147
    CURRENT APPLICATION DATA:
      APPLICATION NUMBER: US/07/493,359
      FILING DATE: 14-MAR-1990
; SEQ ID NO:38:
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RESULT 49
5204326-41
; Patent No. 5204326
    APPLICANT: FUJII, SETSURO; YAMAMOTO, YOSHIHITO; SHIMIZU, FUMIO
; INAI, MASATOSHI; KINOSHITA, NAOSUMI; NAKAMURA, SHIZUO; HIROHASHI,
;MITSURU; SAKAMOTO, TAKASHI;TSUTSUMI, KAZUHIKO;SHIRASAKA, TETSUHIKO
     TITLE OF INVENTION: POLYPEPTIDE DERIVATIVES AND CALCIUM
;METABOLISM IMPROVING AGENT
    NUMBER OF SEQUENCES: 147
     CURRENT APPLICATION DATA:
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;SEQ ID NO:41:
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Query Match
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           5; Conservative 0; Mismatches 0; Indels 0; Gaps
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           6 LQTPQ 10
Db
RESULT 50
5204326-98
; Patent No. 5204326
    APPLICANT: FUJII, SETSURO; YAMAMOTO, YOSHIHITO; SHIMIZU, FUMIO
; INAI, MASATOSHI; KINOSHITA, NAOSUMI; NAKAMURA, SHIZUO; HIROHASHI,
;MITSURU; SAKAMOTO, TAKASHI;TSUTSUMI, KAZUHIKO;SHIRASAKA, TETSUHIKO
    TITLE OF INVENTION: POLYPEPTIDE DERIVATIVES AND CALCIUM
;METABOLISM IMPROVING AGENT
; NUMBER OF SEQUENCES: 147
    CURRENT APPLICATION DATA:
     APPLICATION NUMBER: US/07/493,359
     FILING DATE: 14-MAR-1990
;SEQ ID NO:98:
     LENGTH: 18
5204326-98
 Query Match 29.2%; Score 26; DB 6; Length 18; Best Local Similarity 100.0%; Pred. No. 6.4e+02;
 Matches 5; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
           1 LQTPQ 5
Qу
            6 LQTPQ 10
Db
Search completed: July 4, 2004, 04:48:45
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Job time : 12.6567 secs

## GenCore version 5.1.6 Copyright (c) 1993 - 2004 Compugen Ltd.

OM protein - protein search, using sw model

Run on: July 4, 2004, 04:35:16; Search time 13.0672 Seconds

(without alignments)

125.142 Million cell updates/sec

Title: US-09-641-802-2

Perfect score: 89

Sequence: 1 LQTPQPLLQVMMEPQGD 17

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched: 283366 seqs, 96191526 residues

Total number of hits satisfying chosen parameters: 2898

Minimum DB seq length: 7
Maximum DB seq length: 18

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 100 summaries

Database: PIR 78:\*

1: pir1:\* 2: pir2:\* 3: pir3:\*

4: pir4:\*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

## SUMMARIES

Result No.	Score	% Query Match	Length	DB	ID	Description
1	25	28.1	14	2	C44823	synaptosomal-assoc
2	25	28.1	18	2	A35704	cytochrome P450 ol
3	25	28.1	18	2	F49215	urease (EC 3.5.1.5
4	24	27.0	10	2	A61218	alpha-gliadin 4Ha
5	24	27.0	15	2	A61391	alpha-1-antitrypsi
6	23	25.8	11	2	PC2330	cycloinulooligosac
7	23	25.8	12	2	S70344	amine oxidase (cop
8	23	25.8	15	2	E56819	PS I complex subun
9	23	25.8	16	2	PS0210	28K protein 4209 -
10	23	25.8	16	2	A46236	transforming prote
11	23	25.8	18	2	I59649	human leukocyte an
12	21	23.6	10	2	B61218	alpha-gliadin 6Ha
13	21	23.6	12	2	PH1567	cerebrin 28 - huma

14 15 16 17 18	21 21 21 21 21 21	23.6 23.6 23.6 23.6 23.6 23.6	14 14 15 15 17	2 2 2 2 2 2	PN0147 PH1566 A40634 A42413 PC1318 PN0149
20 21 22	21 20 20 20	23.6 22.5 22.5	18 10 10	2 2 2	S78767 PX0030 PQ0788
23	20	22.5	13	2	D39690
24	20	22.5	14	2	PC7075
25	20	22.5	15	2	PA0021
26	20	22.5	15	2	PN0148
27 28 29	20 20 20	22.5 22.5 22.5 22.5	15 15 16	2 2 2	A41436 PL0110 S00123
30	20	22.5	16	2	D83794
31	20	22.5	17	2	S33609
32	20	22.5	17	2	B31769
33	20	22.5	18	2	S10452
34	20	22.5	18	2	I78841
35	19	21.3	8	2	S21288
36	19	21.3	9	2 2 2	S70345
37	19	21.3	10		D33098
38	19	21.3	10		A61007
39	19	21.3	11	2	S42587
40	19	21.3	11	2	S21127
41	19	21.3	11	2	PN0044
42	19	21.3	12	2	A61309
43	19	21.3	13	2 2 2	S36668
44	19	21.3	14		I54945
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46	19	21.3	14	2	S12904
47	19	21.3	15	2	A56891
48	19	21.3	15	2	F44823
49	19	21.3	16	2 2 2	A24099
50	19	21.3	16		PH0763
51	19	21.3	16		A45454
52	19	21.3	17	2	S41207
53	19	21.3	17	2	S15064
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58	18.5	20.8	14		S00150
59	18.5	20.8	15	2	B41436
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63	18	20.2	11		A34662
64	18	20.2	12		B44818
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69	18	20.2	13	2	A44818
70	18	20.2	13	2	S09716

omega-gliadine 1 a cerebrin 30 - huma orf19 3' of eryK -Ig heavy chain V r large granule L6 c beta-Gliadine 13 ribosomal protein triacylglycerol li NADH2 dehydrogenas neural cell adhesi quanylate cyclase protein QA300016 omega-gliadine 3 alpha-macroglobuli complement factor dihydrolipoamide S hypothetical prote extensin - maize ( T-cell receptor de hypothetical prote thrombopoietin rec lectin - potato (f amine oxidase (cop 214K exoantigen (v hementin (EC 3.4.celF protein - Esc precorrin methyltr protein kinase C i glycoprotein hormo hypothetical prote gene C protein - E actin 8 - slime mo protein kinase (EC gamma 1 gliadin synaptosomal-assoc crystal protein, 2 T-cell receptor be ankyrin-binding gl F420-non-reducinghypothetical prote d(TTAGGG)n-binding u-plasminogen acti gamma2-gliadin P25 kidney stone prote ovostatin - duck ( ovostatin - green brain-associated s exotoxin A - Strep vitellogenin, 190k Achatina cardio-ex extracellular lipa napin - rape (frag napin small chain ATP synthase D cha aminotransferase c extracellular lipa 2S albumin large c

71	18	20.2	13	2	PH1772	T cell receptor al
72	18	20.2	13	2	в47415	mannose-1-phosphat
73	18	20.2	13	2	B56864	dipeptidyl-peptida
74	18	20.2	14	2	PS0278	ribulose-bisphosph
75	18	20.2	15	2	B39109	hypothetical 1.5K
76	18	20.2	15	2	A57003	hypothetical prote
77	18	20.2	15	2	PA0088	protein QF200051 -
78	18	20.2	15	2	A45103	7 alpha-hydroxy-4-
79	18	20.2	15	2	I78838	flt3 ligand isofor
80	18	20.2	16	2	PH0767	T-cell receptor be
81	18	20.2	16	2	PD0002	inulin fructotrans
82	18	20.2	17	2	A60570	Ig mu heavy chain
83	18	20.2	17	2	A35550	adrenocortical cel
84	18	20.2	17	2	B25348	<pre>glycogen(starch) s</pre>
85	18	20.2	18	2	S46418	NTL1 protein - cur
86	18	20.2	18	2	S28408	platelet-derived g
87	18	20.2	18	2	146653	T-cell receptor de
88	18	20.2	18	2	A54195	Na+/K+-exchanging
89	17	19.1	7	2	s71299	ICL2 protein - Par
90	17	19.1	8	2	S53008	citrate synthase -
91	17	19.1	9	2	A41978	calliFMRFamide 1 -
92	17	19.1	9	2	S26508	collagen alpha 2(V
93	17	19.1	9	2	PC7074	translation elonga
94	17	19.1	10	2	S33844	alpha-2-macroglobu
95	17	19.1	10	2	B59272	peptide-N4-(N-acet
96	17	19.1	10	2	S71948	matrix metalloprot
97	17	19.1	10	2	E86128	hypothetical prote
98	17	19.1	11	2	S23306	substance P - Atla
99	17	19.1	11	2	S68637	acetylcholinestera
100	17	19.1	12	2	S10059	tachykinin - Afric

## ALIGNMENTS

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RESULT 1
C44823
synaptosomal-associated protein SNAP-25 peptide 8 - rabbit (fragment)
N; Alternate names: superprotein peptide 8
C; Species: Oryctolagus cuniculus (domestic rabbit)
C; Date: 31-Mar-1993 #sequence revision 18-Nov-1994 #text change 15-Jun-1996
C; Accession: C44823
R; Loewy, A.; Liu, W.S.; Baitinger, C.; Willard, M.B.
J. Neurosci. 11, 3412-3421, 1991
A; Title: The major 35S-methionine-labeled rapidly transported protein
(superprotein) is identical to SNAP-25, a protein of synaptic terminals.
A; Reference number: A44823; MUID: 92044785; PMID: 1941090
A; Accession: C44823
A; Status: preliminary
A; Molecule type: protein
A; Residues: 1-14 <LOE>
A; Experimental source: visual tissue
A; Note: sequence extracted from NCBI backbone (NCBIP:64253)
C; Keywords: membrane trafficking
  Query Match
                          28.1%; Score 25; DB 2; Length 14;
  Best Local Similarity 50.0%; Pred. No. 4.8e+02;
```

```
Matches
             4; Conservative 3; Mismatches 1; Indels 0; Gaps
                                                                              0;
           10 VMMEPQGD 17
Qу
              11:: 11:
            3 VMLDEQGE 10
Db
RESULT 2
A35704
cytochrome P450 olf2 - bovine (fragment)
N; Contains: oxidoreductase (EC 1.-.-.)
C; Species: Bos primigenius taurus (cattle)
C;Date: 31-Mar-1992 #sequence revision 31-Mar-1992 #text change 12-Mar-1999
C; Accession: A35704
R; Lazard, D.; Tal, N.; Rubinstein, M.; Khen, M.; Lancet, D.; Zupko, K.
Biochemistry 29, 7433-7440, 1990
A; Title: Identification and biochemical analysis of novel olfactory-specific
cytochrome P-450IIA and UDP-glucuronosyl transferase.
A; Reference number: A35704; MUID: 91027757; PMID: 2121272
A; Accession: A35704
A; Molecule type: protein
A; Residues: 1-18 <LAZ>
C; Genetics:
A; Gene: CYP2A
C; Superfamily: human cytochrome P450 CYP2D6; cytochrome P450 homology
C; Keywords: electron transfer; heme; monooxygenase; oxidoreductase;
transmembrane protein
                          28.1%; Score 25; DB 2; Length 18;
  Query Match
  Best Local Similarity
                          46.2%; Pred. No. 6.4e+02;
  Matches
            6; Conservative
                                 0; Mismatches
                                                  7; Indels
                                                                 0; Gaps
                                                                              0;
            4 PQPLLQVMMEPQG 16
Qу
              Db
            5 PGPQQQAFKELQG 17
RESULT 3
F49215
urease (EC 3.5.1.5) small chain UreA - Helicobacter mustelae (ATCC 43772)
(fragment)
C; Species: Helicobacter mustelae
C;Date: 19-Dec-1993 #sequence revision 25-Apr-1997 #text change 06-Jan-2003
C; Accession: F49215
R; Turbett, G.R.; Hoj, P.B.; Horne, R.; Mee, B.J.
Infect. Immun. 60, 5259-5266, 1992
A; Title: Purification and characterization of the urease enzymes of Helicobacter
species from humans and animals.
A; Reference number: A49215; MUID: 93084378; PMID: 1452359
A; Accession: F49215
A; Status: preliminary
A; Molecule type: protein
A; Residues: 1-18 <TUR>
A; Note: sequence extracted from NCBI backbone (NCBIP:119487)
C; Superfamily: urease, fused gamma/beta subunit; urease 11K chain homology;
urease 12K chain homology
C; Keywords: hydrolase
```

```
Query Match
                          28.1%; Score 25; DB 2; Length 18;
                          33.3%; Pred. No. 6.4e+02;
  Best Local Similarity
  Matches
             5; Conservative
                                 5; Mismatches
                                                                 0; Gaps
                                                 5; Indels
                                                                             0;
            3 TPQPLLQVMMEPQGD 17
Qy
              11: 1 ::1:
                          |:
            2 TPKELDKMMLHYAGE 16
Db
RESULT 4
A61218
alpha-gliadin 4Ha - grass (Haynaldia villosa) (fragment)
C; Species: Haynaldia villosa, Dasypyrum villosum
C; Date: 19-Mar-1997 #sequence revision 19-Dec-1997 #text change 17-Mar-1999
C; Accession: A61218
R; Shewry, P.R.; Sabelli, P.A.; Parmar, S.; Lafiandra, D.
Biochem. Genet. 29, 207-211, 1991
A; Title: alpha-type prolamins are encoded by genes on chromosomes 4Ha and 6Ha of
Haynaldia villosa Schur (syn. Dasypyrum villosum L.).
A; Reference number: A61218; MUID: 91315394; PMID: 1859356
A; Accession: A61218
A; Status: preliminary
A; Molecule type: protein
A; Residues: 1-10 <SHE>
C; Keywords: seed; storage protein
  Query Match
                          27.0%; Score 24; DB 2; Length 10;
  Best Local Similarity
                          55.6%; Pred. No. 4.9e+02;
  Matches
            5; Conservative
                                1; Mismatches
                                                3; Indels
                                                                 0; Gaps
                                                                             0;
            1 LQTPQPLLQ 9
Qy
              1: | | | |
Db
            2 LRVPVPQLQ 10
RESULT 5
A61391
alpha-1-antitrypsin homolog - human (fragment)
C; Species: Homo sapiens (man)
C;Date: 09-Sep-1994 #sequence revision 09-Sep-1994 #text change 17-Mar-1999
C; Accession: A61391
R; Tanaka, N.; Sekiya, S.; Takamizawa, H.; Kato, N.; Moriyama, Y.; Fujimura, S.
Jpn. J. Cancer Res. 82, 693-700, 1991
A;Title: Characterization of a 54 kDa, alpha-1-antitrypsin-like protein isolated
from ascitic fluid of an endometrial cancer patient.
A; Reference number: A61391; MUID: 91310496; PMID: 1906855
A; Accession: A61391
A; Status: preliminary
A; Molecule type: protein
A; Residues: 1-15 <TAN>
  Query Match
                          27.0%; Score 24; DB 2; Length 15;
  Best Local Similarity
                          100.0%; Pred. No. 7.7e+02;
                                                  0; Indels
  Matches
             4; Conservative
                                0; Mismatches
                                                                 0; Gaps
                                                                             0:
```

Qу

14 PQGD 17

1::||

```
RESULT 6
PC2330
cycloinulooligosaccharide fructanotransferase (EC 2.4.-.-) - Bacillus circulans
(MCI-2554) (fragment)
C; Species: Bacillus circulans
C; Date: 21-Mar-1995 #sequence revision 26-May-1995 #text change 07-May-1999
C; Accession: PC2330
R; Kushibe, S.; Mitsui, K.; Yamagishi, M.; Yamada, K.; Morimoto, Y.
Biosci. Biotechnol. Biochem. 59, 31-34, 1995
A; Title: Purification and characterization of cycloinulooligosaccharide
fructanotransferase (CFTase) from Bacillus circulans MCI-2554.
A; Reference number: PC2330; MUID: 95201377; PMID: 7765973
A; Accession: PC2330
A; Molecule type: protein
A; Residues: 1-11 < KUS>
C; Comment: This enzyme hydrolyzes beta-(2-1) glycosidic linkages and acts in
intermolecular transfructosylation.
C; Keywords: glycosyltransferase; hexosyltransferase
  Query Match
                          25.8%; Score 23; DB 2; Length 11;
                          80.0%; Pred. No. 8e+02;
  Best Local Similarity
                                                                  0; Gaps
 Matches
          . 4; Conservative
                                0; Mismatches
                                                   1; Indels
                                                                              0;
           12 MEPQG 16
Qу
              1 111
Db
            7 MNPOG 11
RESULT 7
S70344
amine oxidase (copper-containing) (EC 1.4.3.6) I - Aspergillus niger (fragments)
C; Species: Aspergillus niger
C; Date: 19-Mar-1998 #sequence revision 17-Apr-1998 #text change 17-Apr-1998
C; Accession: S70344
R; Frebort, I.; Pec, P.; Luhova, L.; Toyama, H.; Matsushita, K.; Hirota, S.;
Kitagawa, T.; Ueno, T.; Asano, Y.; Kato, Y.; Adachi, O.
Biochim. Biophys. Acta 1295, 59-72, 1996
A; Title: Two amine oxidases from Aspergillus niger AKU 3302 contain topa quinone
as the cofactor: unusual cofactor link to the glutamyl residue occurs only at
one of the enzymes.
A; Reference number: S70344; MUID: 96283794; PMID: 8679675
A; Accession: S70344
A; Status: preliminary
A; Molecule type: protein
A; Residues: 1-7;8-12 <FRE>
C; Keywords: oxidoreductase
  Query Match
                          25.8%; Score 23; DB 2; Length 12;
  Best Local Similarity
                          57.1%; Pred. No. 8.8e+02;
                                                                  0; Gaps
             4; Conservative
                                 2; Mismatches
                                                  1; Indels
                                                                              0:
Qу
           10 VMMEPQG 16
```

```
RESULT 8
E56819
PS I complex subunit 8 - cucumber (fragment)
C; Species: Cucumis sativus (cucumber)
C;Date: 05-Jan-1996 #sequence revision 05-Jan-1996 #text_change 05-Jan-1996
C; Accession: E56819
R; Iwasaki, Y.; Ishikawa, H.; Hibino, T.; Takabe, T.
Biochim. Biophys. Acta 1059, 141-148, 1991
A; Title: Characterization of genes that encode subunits of cucumber PS I complex
by N-terminal sequencing.
A; Reference number: A56819; MUID: 91355209; PMID: 1883835
A; Accession: E56819
A; Status: preliminary
A; Molecule type: protein
A; Residues: 1-15 < IWA>
A; Note: sequence extracted from NCBI backbone (NCBIP:58606)
                          25.8%; Score 23; DB 2; Length 15;
  Query Match
  Best Local Similarity
                          30.8%; Pred. No. 1.1e+03;
 Matches
            4; Conservative
                                 3; Mismatches
                                                    6; Indels
                                                                  0; Gaps
            1 LQTPQPLLQVMME 13
Qy
              : | | | | | : :
Db
            1 IQADXPTFQVIQD 13
RESULT 9
PS0210
28K protein 4209 - rice (strain Nihonbare) (fragment)
C; Species: Oryza sativa (rice)
C;Date: 03-Feb-1994 #sequence revision 03-Feb-1994 #text change 11-Apr-1995
C; Accession: PS0210
R;Tsugita, A.; Miyatake, N.
submitted to JIPID, April 1993
A; Reference number: PS0208
A; Accession: PS0210
A; Molecule type: protein
A; Residues: 1-16 <TSA>
A; Experimental source: germ, bran, strain Nihonbare
C; Comment: molecular weight 28K, pI 7.3.
                          25.8%; Score 23; DB 2; Length 16;
  Query Match
                          37.5%; Pred. No. 1.2e+03;
  Best Local Similarity
            3; Conservative 3; Mismatches 2; Indels
                                                                  0; Gaps
           10 VMMEPQGD 17
Qу
              :: 1:11
Db
            8 IVPNPEGD 15
RESULT 10
transforming protein myc, form 1 - mouse (fragment)
C; Species: Mus musculus (house mouse)
```

```
C;Date: 06-Jan-1995 #sequence revision 06-Jan-1995 #text change 06-Jan-1995
C; Accession: A46236
R; Hann, S.R.; Sloan-Brown, K.; Spotts, G.D.
Genes Dev. 6, 1229-1240, 1992
A; Title: Translational activation of the non-AUG-initiated c-myc 1 protein at
high cell densities due to methionine deprivation.
A; Reference number: A46236; MUID: 92331929; PMID: 1628829
A; Accession: A46236
A; Status: preliminary
A; Molecule type: mRNA
A; Residues: 1-16 <HAN>
A; Cross-references: GB:X01023
C: Genetics:
A; Start codon: CTG
C; Keywords: alternative initiators
                          25.8%; Score 23; DB 2; Length 16;
  Query Match
  Best Local Similarity 80.0%; Pred. No. 1.2e+03;
  Matches
             4; Conservative
                               1; Mismatches
                                                 0; Indels
                                                                  0; Gaps
            1 LQTPQ 5
Qу
              1:11
            7 LETPQ 11
Db
RESULT 11
I59649
human leukocyte antigen alpha chain - human (fragment)
C; Species: Homo sapiens (man)
C; Date: 02-Jul-1996 #sequence revision 02-Jul-1996 #text change 05-Nov-1999
C; Accession: I59649
R; Fogdell, A.; Olerup, O.
Tissue Antigens 44, 19-24, 1994
A; Title: The DQA1*0104 allele is carried by DRB1*1001- and DRB1*1401-positive
haplotypes in Caucasians, Africans and Orientals.
A; Reference number: I59649; MUID: 95064785; PMID: 7974465
A; Accession: I59649
A; Status: preliminary; translated from GB/EMBL/DDBJ
A; Molecule type: DNA
A; Residues: 1-18 < RES>
A;Cross-references: GB:S75685; NID:g913771; PIDN:AAB32621.1; PID:g913772
C; Genetics:
A; Gene: GDB: HLA-DQA1
A; Cross-references: GDB:120638; OMIM:146880
A; Map position: 6p21.3-6p21.3
  Query Match
                          25.8%; Score 23; DB 2; Length 18;
  Best Local Similarity 55.6%; Pred. No. 1.4e+03;
  Matches
             5; Conservative
                               0; Mismatches 4; Indels
                                                                  0; Gaps
                                                                              0;
            8 LOVMMEPQG 16
Qу
              1 11 1 1
            5 LTTMMSPCG 13
Db
RESULT 12
```

RESULT 12 B61218

```
alpha-gliadin 6Ha - grass (Haynaldia villosa) (fragment)
C; Species: Haynaldia villosa, Dasypyrum villosum
C;Date: 19-Mar-1997 #sequence revision 19-Dec-1997 #text change 17-Mar-1999
C; Accession: B61218
R; Shewry, P.R.; Sabelli, P.A.; Parmar, S.; Lafiandra, D.
Biochem. Genet. 29, 207-211, 1991
A;Title: alpha-type prolamins are encoded by genes on chromosomes 4Ha and 6Ha of
Haynaldia villosa Schur (syn. Dasypyrum villosum L.).
A; Reference number: A61218; MUID: 91315394; PMID: 1859356
A; Accession: B61218
A; Status: preliminary
A; Molecule type: protein
A; Residues: 1-10 <SHE>
C; Keywords: seed; storage protein
  Query Match
                          23.6%; Score 21; DB 2; Length 10;
  Best Local Similarity
                          44.4%; Pred. No. 1.6e+03;
                                                  3; Indels
  Matches
             4; Conservative
                               2; Mismatches
                                                                  0; Gaps
                                                                              0;
Qу
            1 LQTPQPLLQ 9
              :: | | ||
            1 VRVPVPQLQ 9
Db
RESULT 13
PH1567
cerebrin 28 - human (fragment)
C; Species: Homo sapiens (man)
C; Date: 05-Aug-1994 #sequence revision 05-Aug-1994 #text change 07-May-1999
C; Accession: PH1567
R; Leone, M.G.; Saso, L.; Vecchio, A.D.; Mo, M.; Silvestrini, B.; Cheng, C.Y.
J. Neurochem. 61, 533-540, 1993
A; Title: Micropurification of two human cerebrospinal fluid proteins by high
performance electrophoresis chromatography.
A; Reference number: PH1566; MUID: 93329419; PMID: 8336140
A; Accession: PH1567
A; Molecule type: protein
A; Residues: 1-12 <LEO>
                          23.6%; Score 21; DB 2; Length 12;
  Query Match
  Best Local Similarity
                          44.4%; Pred. No. 1.9e+03;
  Matches
                                2; Mismatches
                                                                              0;
            4; Conservative
                                                   3; Indels
                                                                 0; Gaps
            6 PLLOVMMEP 14
Qу
              | || ::|
Db
            2 PPAQVSVQP 10
RESULT 14
PN0147
omega-gliadine 1 and 2 - Aegilops longissima (fragment)
C; Species: Aegilops longissima
C;Date: 30-Sep-1993 #sequence revision 30-Sep-1993 #text change 07-May-1999
C; Accession: PN0147; PN0146
R;Odintsova, T.I.; Egorov, T.A.
Biokhimiia 55, 509-516, 1990
```

```
A; Title: N-Terminal sequences of omega-gliadins of Aegilops longissima: On the
origin of polyploid wheat genomes.
A; Reference number: PN0146; MUID: 90283493; PMID: 2354218
A; Accession: PN0147
A; Molecule type: protein
A; Residues: 1-14 <ODI>
A; Experimental source: strain K-202
A; Note: omega-gliadine 2 (amino-terminal fragment)
A; Accession: PN0146
A; Molecule type: protein
A; Residues: 1-9 <OD2>
A; Experimental source: strain K-202
A; Note: omega-gliadine 1 (amino-terminal fragment)
  Query Match
                          23.6%; Score 21; DB 2; Length 14;
  Best Local Similarity 100.0%; Pred. No. 2.3e+03;
 Matches
            4; Conservative
                               0; Mismatches
                                                  0; Indels
                                                                  0; Gaps
                                                                              0;
            1 LQTP 4
Qу
              \Box
Db
           11 LQTP 14
RESULT 15
PH1566
cerebrin 30 - human (fragment)
C; Species: Homo sapiens (man)
C;Date: 05-Aug-1994 #sequence revision 05-Aug-1994 #text change 07-May-1999
C; Accession: PH1566
R; Leone, M.G.; Saso, L.; Vecchio, A.D.; Mo, M.; Silvestrini, B.; Cheng, C.Y.
J. Neurochem. 61, 533-540, 1993
A; Title: Micropurification of two human cerebrospinal fluid proteins by high
performance electrophoresis chromatography.
A; Reference number: PH1566; MUID: 93329419; PMID: 8336140
A; Accession: PH1566
A; Molecule type: protein
A; Residues: 1-14 <LEO>
  Query Match
                          23.6%; Score 21; DB 2; Length 14;
  Best Local Similarity 44.4%; Pred. No. 2.3e+03;
  Matches
            4; Conservative 2; Mismatches
                                                  3; Indels
                                                                  0; Gaps
                                                                              0;
            6 PLLQVMMEP 14
Qу
              1 11 ::1
Db
            2 PEAQVSVQP 10
RESULT 16
A40634
orf19 3' of eryK - Saccharopolyspora erythraea (fragment)
C; Species: Saccharopolyspora erythraea
C; Date: 21-Sep-1993 #sequence revision 18-Nov-1994 #text change 18-Nov-1994
C; Accession: A40634
R; Stassi, D.; Donadio, S.; Staver, M.J.; Katz, L.
J. Bacteriol. 175, 182-189, 1993
A; Title: Identification of a Saccharopolyspora erythraea gene required for the
final hydroxylation step in erythromycin biosynthesis.
```

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A; Reference number: A40634; MUID: 93106953; PMID: 8416893
A; Accession: A40634
A; Status: preliminary
A; Molecule type: nucleic acid
A; Residues: 1-15 <STA>
A; Note: sequence extracted from NCBI backbone (NCBIN:121243, NCBIP:121244)
  Query Match
                          23.6%; Score 21; DB 2; Length 15;
  Best Local Similarity
                          50.0%; Pred. No. 2.5e+03;
                                                  1; Indels
 Matches
             3; Conservative
                                 2; Mismatches
                                                                  0; Gaps
                                                                              0:
            1 LQTPQP 6
Qy
              : ||:|
Db
            7 VSTPRP 12
RESULT 17
A42413
Ig heavy chain V region (LBL-copurifying) - rabbit (fragment)
C; Species: Oryctolagus cuniculus (domestic rabbit)
C;Date: 31-Dec-1993 #sequence revision 31-Dec-1993 #text change 07-May-1999
C; Accession: A42413
R; Bao, Z.; Muschler, J.; Horwitz, A.F.
J. Biol. Chem. 267, 4974-4980, 1992
A; Title: LBL, a novel, developmentally regulated, laminin-binding lectin.
A; Reference number: A42413; MUID: 92165867; PMID: 1531660
A; Accession: A42413
A; Status: preliminary
A; Molecule type: protein
A; Residues: 1-15 <BAO>
C; Keywords: heterotetramer; immunoglobulin
  Query Match
                          23.6%; Score 21; DB 2; Length 15;
  Best Local Similarity 100.0%; Pred. No. 2.5e+03;
             4; Conservative
                                0; Mismatches
                                                   0; Indels
                                                                  0; Gaps
                                                                              0;
            1 LQTP 4
Qу
              1111
Db
           11 LQTP 14
RESULT 18
PC1318
large granule L6 chain - horseshoe crab (Tachypleus tridentatus) (fragment)
C; Species: Tachypleus tridentatus
C; Date: 10-Mar-1994 #sequence revision 10-Mar-1994 #text change 07-May-1999
C; Accession: PC1318
R; Shigenaga, T.; Takayenoki, Y.; Kawasaki, S.; Seki, N.; Muta, T.; Toh, Y.; Ito,
A.; Iwanaga, S.
J. Biochem. 114, 307-316, 1993
A; Title: Separation of large and small granules from horseshoe crab (Tachypleus
tridentatus) hemocytes and characterization of their components.
A; Reference number: PC1309; MUID: 94110249; PMID: 8282718
A; Accession: PC1318
A; Molecule type: protein
A; Residues: 1-17 <SHI>
C; Comment: This protein participates in immobilization of invading microbes.
```

```
Query Match
                          23.6%; Score 21; DB 2; Length 17;
  Best Local Similarity
                          30.8%; Pred. No. 2.8e+03;
                                                                 0; Gaps
             4; Conservative
                                 2; Mismatches
                                                 7; Indels
                                                                             0;
            2 QTPQPLLQVMMEP 14
Qу
              1 | |::
                         - 1
            5 QIPGKLMHITATP 17
Db
RESULT 19
PN0149
beta-Gliadine 13 - Aegilops longissima (fragment)
C; Species: Aegilops longissima
C;Date: 30-Sep-1993 #sequence revision 30-Sep-1993 #text change 07-May-1999
C; Accession: PN0149
R;Odintsova, T.I.; Egorov, T.A.
Biokhimiia 55, 509-516, 1990
A; Title: N-Terminal sequences of omega-gliadins of Aegilops longissima: On the
origin of polyploid wheat genomes.
A; Reference number: PN0146; MUID: 90283493; PMID: 2354218
A; Accession: PN0149
A; Molecule type: protein
A; Residues: 1-18 <ODI>
A; Experimental source: strain K-202
C; Superfamily: gliadin
  Query Match
                          23.6%; Score 21; DB 2; Length 18;
                          44.4%; Pred. No. 3e+03;
  Best Local Similarity
  Matches
            4; Conservative
                                2; Mismatches
                                                  3; Indels
                                                                 0; Gaps
                                                                             0;
            1 LOTPOPLLO 9
Qy
              :: | | | |
            1 VRVPVPQLQ 9
Db
RESULT 20
S78767
ribosomal protein MRP-S29, mitochondrial - bovine (fragment)
C; Species: Bos primigenius taurus (cattle)
C; Date: 20-Sep-1999 #sequence revision 20-Sep-1999 #text change 20-Sep-1999
C; Accession: S78767
R; Graack, H.R.
submitted to the Protein Sequence Database, July 1999
A; Reference number: S78760
A; Accession: S78767
A; Molecule type: protein
A; Residues: 1-18 <GRA>
C; Keywords: mitochondrion
F;1-18/Product: ribosomal protein MRP-S29 (fragment) #status experimental <MAT>
  Query Match
                          23.6%; Score 21; DB 2; Length 18;
  Best Local Similarity
                          33.3%; Pred. No. 3e+03;
  Matches
             3; Conservative
                                3; Mismatches
                                                  3; Indels
                                                                 0; Gaps
                                                                             0;
            7 LLQVMMEPQ 15
Qу
              :1: 1 1:
```

```
RESULT 21
PX0030
triacylglycerol lipase (EC 3.1.1.3) II - yeast (Geotrichum candidum) (fragments)
C; Species: Geotrichum candidum
C;Date: 31-Dec-1990 #sequence revision 31-Dec-1990 #text change 30-Sep-1993
C; Accession: PX0030
R; Sugihara, A.; Shimada, Y.; Tominaga, Y.
J. Biochem. 107, 426-430, 1990
A; Title: Separation and characterization of two molecular forms of Geotrichum
candidum lipase.
A; Reference number: PX0030; MUID: 90256718; PMID: 2341377
A; Accession: PX0030
A; Molecule type: protein
A; Residues: 1-10 <SUG>
C; Comment: Lipase catalyzes the hydrolysis of briacylglycerols. This fungus
contains two forms of lipase, lipase I and lipase II.
C; Keywords: carboxylic ester hydrolase
  Query Match
                          22.5%; Score 20; DB 2; Length 10;
  Best Local Similarity
                          42.9%; Pred. No. 2.3e+03;
             3; Conservative
                                 2; Mismatches
                                                  2; Indels
                                                                  0; Gaps
                                                                              0;
            2 QTPQPLL 8
Qу
              : |:| |
Db
            1 EAPRPXL 7
RESULT 22
PO0788
NADH2 dehydrogenase (EC 1.6.99.3) 20K chain - fava bean mitochondrion (fragment)
N; Alternate names: complex I 20K chain; NADH-ubiquinone reductase 20K chain
C; Species: mitochondrion Vicia faba (fava bean)
C; Date: 03-May-1994 #sequence revision 07-Oct-1994 #text change 03-Jun-2002
C; Accession: PQ0788
R; Leterme, S.; Boutry, M.
Plant Physiol. 102, 435-443, 1993
A; Title: Purification and preliminary characterization of mitochondrial complex
I (NADH: ubiquinone reductase) from broad bean (Vicia faba L.).
A; Reference number: PQ0775; MUID: 94151437; PMID: 8108509
A; Accession: PQ0788
A; Molecule type: protein
A; Residues: 1-10 <LET>
C; Comment: Complex I, mitochondrial NADH-ubiquiquinone reductase, is the first
of the three proton-translocating complexes of the mitochondrial respiratory
chain and composed of 35 different subunits ranging from 5K to 75K.
C; Comment: This enzyme catalyzes electron transfer from endogenous NADH to
ubiquinone by a large number of redox groups.
C; Genetics:
A; Genome: mitochondrion
C; Keywords: electron transfer; mitochondrion; oxidoreductase
  Query Match
                          22.5%; Score 20; DB 2; Length 10;
  Best Local Similarity
                          83.3%; Pred. No. 2.3e+03;
  Matches
             5; Conservative
                                 0; Mismatches
                                                 1; Indels 0; Gaps
```

```
5 QPLLQV 10
Qу
              1 1111
            5 QVLLQV 10
Db
RESULT 23
D39690
neural cell adhesion molecule, cardiac splice form +,-,-,+ - rat (fragment)
C; Species: Rattus norvegicus (Norway rat)
C;Date: 24-Jan-1992 #sequence revision 24-Jan-1992 #text change 16-Jul-1999
C; Accession: D39690
R; Reyes, A.A.; Small, S.J.; Akeson, R.
Mol. Cell. Biol. 11, 1654-1661, 1991
A; Title: At least 27 alternatively spliced forms of the neural cell adhesion
molecule mRNA are expressed during rat heart development.
A; Reference number: A39690; MUID: 91141516; PMID: 1996115
A; Accession: D39690
A; Status: preliminary; nucleic acid sequence not shown; not compared with
conceptual translation
A; Molecule type: mRNA
A; Residues: 1-13 < REY>
A; Cross-references: GB:M63970
C; Superfamily: neural cell adhesion molecule; fibronectin type III repeat
homology; immunoglobulin homology
C; Keywords: cardiac muscle; cell adhesion; heart
  Query Match
                          22.5%; Score 20; DB 2; Length 13;
                          75.0%; Pred. No. 3.1e+03;
  Best Local Similarity
  Matches
            3; Conservative 1; Mismatches 0; Indels
                                                                 0; Gaps
                                                                             0;
           14 POGD 17
Qу
              111:
Db
            8 PQGE 11
RESULT 24
PC7075
guanylate cyclase (EC 4.6.1.2), soluble, alpha-1 chain - mouse (fragment)
C; Species: Mus musculus (house mouse)
C;Date: 18-Aug-2000 #sequence revision 18-Aug-2000 #text change 18-Aug-2000
C; Accession: PC7075
R;Tsugita, A.; Kawakami, T.; Uchida, T.; Sakai, T.; Kamo, M.; Matsui, T.;
Watanabe, Y.; Morimasa, T.; Hosokawa, K.; Toda, T.
Electrophoresis 21, 1853-1871, 2000
A; Title: Proteome analysis of mouse brain: Two-dimensional electrophoresis
profiles of tissue proteins during the course of aging.
A; Reference number: PC7072
A; Accession: PC7075
A; Molecule type: protein
A; Residues: 1-14 <TSU>
A; Experimental source: strain C57BL/6Cr Slc, male; brain, striatum
C; Keywords: brain; phosphorus-oxygen lyase
  Query Match
                          22.5%; Score 20; DB 2; Length 14;
  Best Local Similarity
                          75.0%; Pred. No. 3.3e+03;
             3; Conservative 1; Mismatches
  Matches
                                                  0; Indels
                                                                 0; Gaps
```

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14 PQGD 17
Qу
              111:
Db
            9 PQGE 12
RESULT 25
PA0021
protein QA300016 - Arabidopsis thaliana (fragment)
C; Species: Arabidopsis thaliana (mouse-ear cress)
C;Date: 30-Jun-1992 #sequence revision 06-Jan-1995 #text change 06-Jun-1997
C; Accession: PA0021
R; Kamo, M.; Kawakami, T.; Miyatake, N.; Tsugita, A.
submitted to JIPID, July 1994
A; Description: Separation and characterization of Arabidopsis proteins by two-
dimensional gel electrophoresis.
A; Reference number: PA0001
A; Accession: PA0021
A; Molecule type: protein
A; Residues: 1-15 < KAM>
A; Experimental source: leaf
  Query Match
                          22.5%; Score 20; DB 2; Length 15;
  Best Local Similarity
                          42.9%; Pred. No. 3.6e+03;
                                                                  0; Gaps
  Matches
             3; Conservative
                                 3; Mismatches
                                                   1; Indels
                                                                               0;
Qу
            9 OVMMEPO 15
              :::!!!
Db
            5 EXVIEPQ 11
RESULT 26
PN0148
omega-gliadine 3 - Aegilops longissima (fragment)
C; Species: Aegilops longissima
C;Date: 30-Sep-1993 #sequence revision 30-Sep-1993 #text change 07-May-1999
C; Accession: PN0148
R;Odintsova, T.I.; Egorov, T.A.
Biokhimiia 55, 509-516, 1990
A; Title: N-Terminal sequences of omega-gliadins of Aegilops longissima: On the
origin of polyploid wheat genomes.
A; Reference number: PN0146; MUID: 90283493; PMID: 2354218
A; Accession: PN0148
A; Molecule type: protein
A; Residues: 1-15 <ODI>
A; Experimental source: strain K-202
A; Note: 11-Gln was also found
  Query Match
                          22.5%; Score 20; DB 2; Length 15;
                                  Pred. No. 3.6e+03;
  Best Local Similarity
                          80.0%;
                                                 1; Indels
  Matches
             4; Conservative
                                 0; Mismatches
                                                                  0; Gaps
                                                                               0;
Qy
            1 LQTPQ 5
              1 + 1 + 1
Db
           11 LXTPQ 15
```

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RESULT 27
A41436
alpha-macroglobulin - green seaturtle (fragment)
C; Species: Chelonia mydas (green seaturtle)
C; Date: 21-Apr-1992 #sequence revision 21-Apr-1992 #text change 18-Jun-1993
C; Accession: A41436
R;Osada, T.; Sasaki, T.; Ikai, A.
J. Biochem. 103, 212-217, 1988
A; Title: Purification and characterization of alpha-macroglobulin and
ovomacroglobulin of the green turtle (Chelonia mydas japonica).
A; Reference number: A41436; MUID: 88227890; PMID: 2453503
A; Accession: A41436
A; Status: preliminary
A; Molecule type: protein
A; Residues: 1-15 <OSA>
  Query Match
                          22.5%; Score 20; DB 2; Length 15;
  Best Local Similarity 33.3%; Pred. No. 3.6e+03;
  Matches
             3; Conservative 3; Mismatches 3; Indels
                                                                 0; Gaps
                                                                              0;
            4 PQPLLQVMM 12
Qу
             |:| |::
Db
            2 PEPQYMVLV 10
RESULT 28
PL0110
complement factor B1-Bb and B2-Bb - guinea pig (fragment)
C; Species: Cavia porcellus (guinea pig)
C;Date: 31-Mar-1990 #sequence revision 31-Mar-1990 #text change 15-Nov-1996
C; Accession: PL0110
R; Matsushita, M.; Okada, H.
Mol. Immunol. 26, 669-676, 1989
A; Title: Two forms of guinea pig factor B of the alternative complement pathway
with different molecular weights.
A; Reference number: A93136; MUID: 89384686; PMID: 2779589
A; Accession: PL0110
A; Molecule type: protein
A; Residues: 1-15 <MAT>
C; Keywords: complement alternate pathway; glycoprotein
                          22.5%; Score 20; DB 2; Length 15;
  Query Match
                          28.6%; Pred. No. 3.6e+03;
  Best Local Similarity
  Matches
             2; Conservative
                                 4; Mismatches
                                                                              0;
                                                  1; Indels 0; Gaps
           10 VMMEPQG 16
Qу
              ::::| |
Db
            2 IILDPAG 8
RESULT 29
S00123
dihydrolipoamide S-succinyltransferase (EC 2.3.1.61) - bovine (fragment)
N; Alternate names: 2-oxoglutarate dehydrogenase complex chain E2;
transsuccinylase
C; Species: Bos primigenius taurus (cattle)
C;Date: 07-Sep-1990 #sequence_revision 07-Sep-1990 #text_change 05-May-2000
```

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C; Accession: S00123
R; Bradford, A.P.; Aitken, A.; Beg, F.; Cook, K.G.; Yeaman, S.J.
FEBS Lett. 222, 211-214, 1987
A; Title: Amino acid sequence surrounding the lipoic acid cofactor of bovine
kidney 2-oxoglutarate dehydrogenase complex.
A; Reference number: S00123; MUID: 88005143; PMID: 3115829
A; Accession: S00123
A; Molecule type: protein
A; Residues: 1-16 <BRA>
C; Superfamily: dihydrolipoamide acetyltransferase; lipoyl/biotin-binding
homology
C; Keywords: acyltransferase; coenzyme A; lipoamide; mitochondrion;
oxidoreductase; tricarboxylic acid cycle
F;1-16/Domain: lipoyl/biotin-binding homology (fragment) <LPB>
F;5/Binding site: lipoamide (Lys) (covalent) #status experimental
                          22.5%; Score 20; DB 2; Length 16;
  Query Match
  Best Local Similarity 50.0%; Pred. No. 3.9e+03;
 Matches
             3; Conservative
                              1; Mismatches 2; Indels
                                                                 0; Gaps
                                                                             0;
            1 LQTPQP 6
Qу
             : 1 1 1
Db
            8 VQVPSP 13
RESULT 30
D83794
hypothetical protein BH1156 [imported] - Bacillus halodurans (strain C-125)
C; Species: Bacillus halodurans
C;Date: 01-Dec-2000 #sequence revision 01-Dec-2000 #text change 15-Jun-2001
C; Accession: D83794
R; Takami, H.; Nakasone, K.; Takaki, Y.; Maeno, G.; Sasaki, R.; Masui, N.; Fuji,
F.; Hirama, C.; Nakamura, Y.; Ogasawara, N.; Kuhara, S.; Horikoshi, K.
Nucleic Acids Res. 28, 4317-4331, 2000
A; Title: Complete genome sequence of the alkaliphilic bacterium Bacillus
halodurans and genomic sequence comparison with Bacillus subtilis.
A; Reference number: A83650; MUID:20512582; PMID:11058132
A; Accession: D83794
A; Status: preliminary
A; Molecule type: DNA
A; Residues: 1-16 <STO>
A;Cross-references: GB:AP001511; GB:BA000004; NID:g10173727; PIDN:BAB04875.1;
GSPDB:GN00137
A; Experimental source: strain C-125
C; Genetics:
A; Gene: BH1156
  Query Match
                          22.5%; Score 20; DB 2; Length 16;
  Best Local Similarity 66.7%; Pred. No. 3.9e+03;
  Matches
             4; Conservative
                              1; Mismatches
                                                  1; Indels
                                                                 0; Gaps
                                                                             0;
           11 MMEPQG 16
Qу
              111:1
Db
           10 MMELEG 15
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s33609
extensin - maize (fragment)
C; Species: Zea mays (maize)
C;Date: 19-Mar-1997 #sequence revision 11-Jun-1999 #text_change 11-Jun-1999
C; Accession: S33609
R; Murphy, J.M.; Hood, E.E.
Plant Mol. Biol. 21, 885-893, 1993
A; Title: Molecular basis for extensin size heterogeneity in two maize varieties.
A; Reference number: S33609; MUID: 93222485; PMID: 8467081
A; Accession: S33609
A; Molecule type: protein
A; Residues: 1-17 < MUR>
C; Keywords: glycoprotein; hydroxyproline
  Query Match
                          22.5%; Score 20; DB 2; Length 17;
  Best Local Similarity 75.0%; Pred. No. 4.2e+03;
  Matches
             3; Conservative 1; Mismatches 0; Indels
                                                                 0; Gaps
                                                                              0;
            3 TPQP 6
Qy
              11:1
Db
           13 TPKP 16
RESULT 32
B31769
T-cell receptor delta-2 chain J region - human (fragment)
C; Species: Homo sapiens (man)
C;Date: 07-Jun-1990 #sequence revision 07-Jun-1990 #text change 05-Nov-1999
C; Accession: B31769
R; Loh, E.Y.; Cwirla, S.; Serafini, A.T.; Phillips, J.H.; Lanier, L.L.
Proc. Natl. Acad. Sci. U.S.A. 85, 9714-9718, 1988
A; Title: Human T-cell-receptor delta chain: genomic organization, diversity, and
expression in populations of cells.
A; Reference number: A94221; MUID: 89071766; PMID: 2974163
A; Accession: B31769
A; Molecule type: DNA
A; Residues: 1-17 <LOH>
A;Cross-references: GB:L36386; NID:g540455; PIDN:AAA61108.1; PID:g540456
C; Keywords: T-cell receptor
                          22.5%; Score 20; DB 2; Length 17;
  Query Match
                          50.0%; Pred. No. 4.2e+03;
  Best Local Similarity
             3; Conservative
                                3; Mismatches
                                                0; Indels
                                                                 0; Gaps
                                                                              0;
            9 QVMMEP 14
Qу
              1:::11
Db
           12 QLIVEP 17
RESULT 33
S10452
hypothetical protein (bphA 5' region) - Aspergillus niger
C; Species: Aspergillus niger
C;Date: 31-Dec-1990 #sequence revision 31-Dec-1990 #text change 09-Sep-1997
C; Accession: S10452
R; van Gorcom, R.F.M.; Boschloo, J.G.; Kuijvenhoven, A.; Lange, J.; Bos, C.J.;
van Balken, J.A.M.; Pouwels, P.H.; van den Hondel, C.A.M.J.J.
```

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submitted to the EMBL Data Library, March 1990
A; Reference number: S10452
A; Accession: S10452
A; Molecule type: DNA
A; Residues: 1-18 < VAN>
A; Cross-references: EMBL: X52521; NID: g2336; PID: g2337
                          22.5%; Score 20; DB 2; Length 18;
  Query Match
  Best Local Similarity
                          80.0%; Pred. No. 4.4e+03;
  Matches
            4; Conservative
                                0; Mismatches
                                                  1; Indels
                                                                  0; Gaps
                                                                               0:
            3 TPOPL 7
Qу
              1 111
Db
            2 TDQPL 6
RESULT 34
I78841
thrombopoietin receptor - mouse (fragment)
C; Species: Mus sp. (mouse)
C;Date: 02-Aug-1996 #sequence revision 02-Aug-1996 #text change 05-Nov-1999
C; Accession: I78841
R; Alexander, W.S.; Dunn, A.R.
Oncogene 10, 795-803, 1995
A; Title: Structure and transcription of the genomic locus encoding murine c-Mpl,
a receptor for thrombopoietin.
A; Reference number: I58350; MUID: 95166571; PMID: 7862460
A; Accession: I78841
A; Status: preliminary; translated from GB/EMBL/DDBJ
A; Molecule type: mRNA
A; Residues: 1-18 < RES>
A;Cross-references: GB:S76842; NID:q912990; PIDN:AAB33462.1; PID:q912991
C; Genetics:
A; Gene: c-mpl1
                          22.5%; Score 20; DB 2; Length 18;
  Query Match
  Best Local Similarity 57.1%; Pred. No. 4.4e+03;
            4; Conservative 0; Mismatches 3; Indels
  Matches
                                                                  0; Gaps
                                                                              0;
Qу
            4 PQPLLQV 10
              I I I I
            5 PAPFLTV 11
Db
RESULT 35
S21288
lectin - potato (fragment)
C; Species: Solanum tuberosum (potato)
C;Date: 19-Mar-1997 #sequence_revision 05-Dec-1998 #text_change 05-Dec-1998
C; Accession: S21288
R; Millar, D.J.; Allen, A.K.; Smith, C.G.; Sidebottom, C.; Slabas, A.R.; Bolwell,
G.P.
Biochem. J. 283, 813-821, 1992
A; Title: Chitin-binding proteins in potato (Solanum tuberosum L.) tuber.
Characterization, immunolocalization and effects of wounding.
A; Reference number: S21288; MUID: 92272683; PMID: 1590771
A; Accession: S21288
```

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A; Molecule type: protein
A; Residues: 1-8 <MIL>
A; Experimental source: var. Ulster Sceptre
A; Description: may be involved in defence mechanism of the plant
C; Keywords: hydroxyproline; lectin
  Query Match
                          21.3%; Score 19; DB 2; Length 8;
  Best Local Similarity
                          75.0%; Pred. No. 2.8e+05;
  Matches
            3; Conservative
                                0; Mismatches
                                                  1; Indels
                                                                0; Gaps
                                                                              0;
            3 TPOP 6
Qy
              \perp
Db
            4 TPSP 7
RESULT 36
S70345
amine oxidase (copper-containing) (EC 1.4.3.6) II - Aspergillus niger
(fragments)
C; Species: Aspergillus niger
C;Date: 19-Mar-1998 #sequence_revision 17-Apr-1998 #text_change 17-Apr-1998
C; Accession: S70345
R; Frebort, I.; Pec, P.; Luhova, L.; Toyama, H.; Matsushita, K.; Hirota, S.;
Kitagawa, T.; Ueno, T.; Asano, Y.; Kato, Y.; Adachi, O.
Biochim. Biophys. Acta 1295, 59-72, 1996
A; Title: Two amine oxidases from Aspergillus niger AKU 3302 contain topa quinone
as the cofactor: unusual cofactor link to the glutamyl residue occurs only at
one of the enzymes.
A; Reference number: S70344; MUID: 96283794; PMID: 8679675
A; Accession: S70345
A; Status: preliminary
A; Molecule type: protein
A; Residues: 1-5;6-9 <FRE>
C; Keywords: oxidoreductase
  Query Match
                          21.3%; Score 19; DB 2; Length 9;
  Best Local Similarity 37.5%; Pred. No. 2.8e+05;
             3; Conservative
                                3; Mismatches
                                                  2; Indels
                                                                  0; Gaps
                                                                              0;
Qу
           10 VMMEPQGD 17
              1::11
Db
            1 VVIEPNXE 8
RESULT 37
D33098
214K exoantigen (version 1) - malaria parasite (Plasmodium falciparum)
(fragments)
C; Species: Plasmodium falciparum
C;Date: 24-Aug-1990 #sequence revision 24-Aug-1990 #text change 09-Jun-2000
C; Accession: D33098
R; Nichols, J.H.; Hager, L.P.
submitted to the Protein Sequence Database, May 1990
A; Reference number: A33098
A; Accession: D33098
A; Status: preliminary
```

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A; Molecule type: protein
A; Residues: 1-10 <NIC>
  Query Match
                          21.3%; Score 19; DB 2; Length 10;
  Best Local Similarity 75.0%; Pred. No. 3.4e+03;
  Matches
             3; Conservative 1; Mismatches
                                                 0; Indels
                                                                 0; Gaps
           11 MMEP 14
Qу
             1:11
Db
            4 MLEP 7
RESULT 38
A61007
hementin (EC 3.4.-.-) - Amazon leech (fragment)
C; Species: Haementeria ghilianii (Amazon leech)
C;Date: 31-Dec-1993 #sequence revision 31-Dec-1993 #text change 07-May-1999
C; Accession: A61007
R; Swadesh, J.K.; Huang, I.Y.; Budzynski, A.Z.
J. Chromatogr. 502, 359-369, 1990
A; Title: Purification and characterization of hementin, a fibrinogenolytic
protease from the leech Haementeria ghilianii.
A; Reference number: A61007; MUID: 90256973; PMID: 2187898
A; Accession: A61007
A; Molecule type: protein
A; Residues: 1-10 <SWA>
C; Keywords: anticoagulant; hydrolase; saliva
  Query Match
                          21.3%; Score 19; DB 2; Length 10;
                          42.9%; Pred. No. 3.4e+03;
  Best Local Similarity
           3; Conservative
  Matches
                                2: Mismatches
                                                2; Indels
                                                                0; Gaps
           11 MMEPQGD 17
Qу
              : ||: |
            3 LTEPEPD 9
Db
RESULT 39
S42587
celF protein - Escherichia coli (fragment)
C; Species: Escherichia coli
C;Date: 25-Dec-1994 #sequence revision 10-Nov-1995 #text change 10-Nov-1995
C; Accession: S42587
R; Guzzo, A.; DuBow, M.S.
Mol. Gen. Genet. 242, 455-460, 1994
A; Title: A luxAB transcriptional fusion to the cryptic celf gene of Escherichia
coli displays increased luminescence in the presence of nickel.
A; Reference number: S42587; MUID: 94166755; PMID: 8121401
A; Accession: S42587
A; Molecule type: DNA
A; Residues: 1-11 <GUZ>
C; Genetics:
A; Gene: celF
                          21.3%; Score 19; DB 2; Length 11;
  Query Match
                          75.0%; Pred. No. 3.7e+03;
  Best Local Similarity
  Matches
            3; Conservative
                               0; Mismatches
                                                 1; Indels 0; Gaps
```

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3 TPQP 6
Qу
             -1111
            6 TPHP 9
Db
RESULT 40
S21127
precorrin methyltransferase - Salmonella typhimurium
C; Species: Salmonella typhimurium
C; Date: 22-Nov-1993 #sequence revision 10-Nov-1995 #text change 10-Nov-1995
C; Accession: S21127
R; Roessper, C.A.; Warren, M.J.; Santander, P.J.; Atshaves, B.P.; Ozaki, S.;
Stolowich, N.J.; Iida, K.; Scott, A.I.
FEBS Lett. 301, 73-78, 1992
A; Title: Expression of 9 Salmonella typhimurium enzymes for cobinamide
synthesis. Identification of the 11-methyl and 20-methyl transferases of corrin
biosynthesis.
A; Reference number: S21127; MUID: 93083628; PMID: 1451790
A; Accession: S21127
A;Status: preliminary
A; Molecule type: protein
A; Residues: 1-11 < ROE>
  Query Match
                          21.3%; Score 19; DB 2; Length 11;
  Best Local Similarity
                          42.9%; Pred. No. 3.7e+03;
 Matches
             3; Conservative
                                2; Mismatches
                                                   2; Indels
                                                                  0; Gaps
                                                                              0;
            1 LOTPOPL 7
Qу
              : | | | :
Db
            4 IQOPOTI 10
RESULT 41
PN0044
protein kinase C inhibitor I - mouse (fragment)
C; Species: Mus musculus (house mouse)
C; Date: 29-Oct-1997 #sequence revision 29-Oct-1997 #text change 23-Jan-1998
C; Accession: PN0044
R; Kato, H.
Kawasaki Igakkaishi 22, 245-259, 1996
A; Title: Analysis of proteins isolated by two dimensional electrophoresis of
mouse neuroblastoma cells.
A: Reference number: PN0041
A; Accession: PN0044
A; Molecule type: protein
A; Residues: 1-11 <KAT>
A; Experimental source: neuroblastoma cell
C; Comment: The molecular mass is 13,900 and the pI is 6.36. The amino-terminus
is blocked.
C; Keywords: brain
  Query Match
                          21.3%; Score 19; DB 2; Length 11;
  Best Local Similarity
                          60.0%; Pred. No. 3.7e+03;
                                                                  0; Gaps
  Matches
             3: Conservative
                                1; Mismatches
                                                   1; Indels
                                                                              0;
Qу
           13 EPQGD 17
```

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RESULT 42
A61309
glycoprotein hormones alpha chain - hamster (fragment)
N; Alternate names: choriogonadotropin alpha chain; luteinizing hormone alpha
chain; lutropin alpha chain; thyrotropin alpha chain
C; Species: Cricetinae gen. sp. (hamster)
C; Date: 17-Jul-1994 #sequence revision 17-Jul-1994 #text change 07-May-1999
C; Accession: A61309
R; Glenn, S.D.; Nahm, H.S.; Greenwald, G.S.; Ward, D.N.
Endocrinology 111, 1263-1269, 1982
A; Title: Isolation and characterization of hamster luteinizing hormone.
A; Reference number: A61309; MUID: 83003498; PMID: 6889489
A; Accession: A61309
A; Status: preliminary
A; Molecule type: protein
A; Residues: 1-12 <GLE>
C; Superfamily: glycoprotein hormones alpha chain
C; Keywords: glycoprotein
  Query Match
                          21.3%; Score 19; DB 2; Length 12;
  Best Local Similarity
                          75.0%; Pred. No. 4.1e+03;
             3; Conservative
                                 0; Mismatches
                                                   1; Indels
                                                                  0; Gaps
                                                                              0;
           14 PQGD 17
Qу
              1 11
Db
            2 PBGD 5
RESULT 43
S36668
hypothetical protein 133 - Plectonema boryanum (fragment)
C; Species: Plectonema boryanum
C;Date: 31-Dec-1993 #sequence revision 31-Dec-1993 #text change 15-Oct-1999
C; Accession: S36668
R; Fujita, Y.; Matsumoto, H.; Takahashi, Y.; Matsubara, H.
submitted to the EMBL Data Library, August 1992
A; Description: Identification of the nifDK-like gene (ORF467) involved in the
biosynthesis of chlorophyll in the cyanobacterium Plectonema boryanum.
A; Reference number: S36668
A; Accession: S36668
A; Status: preliminary
A; Molecule type: DNA
A; Residues: 1-13 <FUJ>
A; Cross-references: EMBL: D12973; NID: q216810; PIDN: BAA02348.1; PID: d1002847;
PID:g216811
  Query Match
                          21.3%; Score 19; DB 2; Length 13;
  Best Local Similarity
                          75.0%; Pred. No. 4.5e+03;
  Matches
             3; Conservative
                                0; Mismatches
                                                  1; Indels
                                                                  0; Gaps
                                                                              0;
            3 TPQP 6
Qу
              \Box
Db
            9 TPSP 12
```

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RESULT 44
154945
gene C protein - Escherichia coli (fragment)
C; Species: Escherichia coli
C;Date: 07-Jun-1996 #sequence revision 07-Jun-1996 #text change 08-Oct-1999
C; Accession: I54945
R; Tao, T.; Bourne, J.C.; Blumenthal, R.M.
J. Bacteriol. 173, 1367-1375, 1991
A; Title: A family of regulatory genes associated with type II restriction-
modification systems.
A; Reference number: I54945; MUID: 91139577; PMID: 1995588
A; Accession: I54945
A; Status: preliminary; translated from GB/EMBL/DDBJ
A; Molecule type: DNA
A; Residues: 1-14 < RES>
A;Cross-references: GB:M63619; NID:g147664; PIDN:AAA24555.1; PID:g147665
  Query Match
                          21.3%; Score 19; DB 2; Length 14;
                          57.1%; Pred. No. 4.9e+03;
  Best Local Similarity
            4; Conservative
  Matches
                                 2; Mismatches
                                                 1; Indels
                                                                 0; Gaps
                                                                              0;
Qу
            7 LLQVMME 13
              1:1:11
            1 LKEVIME 7
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A39239
actin 8 - slime mold (Dictyostelium discoideum) (fragment)
C; Species: Dictyostelium discoideum
C;Date: 20-Sep-1991 #sequence revision 20-Sep-1991 #text change 18-Jun-1993
C; Accession: A39239
R; Frankel, S.; Condeelis, J.; Leinwand, L.
J. Biol. Chem. 265, 17980-17987, 1990
A; Title: Expression of actin in Escherichia coli. Aggregation, solubilization,
and functional analysis.
A; Reference number: A39239; MUID: 91009269; PMID: 2211676
A; Accession: A39239
A; Status: preliminary
A; Molecule type: protein
A; Residues: 1-14 <FRA>
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protein kinase (EC 2.7.1.37) - starfish (Pisaster ochraceus)
C; Species: Pisaster ochraceus
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C; Date: 19-Mar-1997 #sequence revision 19-Mar-1997 #text change 01-Aug-1997
C; Accession: S12904
R; Sanghera, J.S.; Aebersold, R.; Morrison, H.D.; Bures, E.J.; Pelech, S.L.
FEBS Lett. 273, 223-226, 1990
A; Title: Identification of the sites in myelin basic protein that are
phosphorylated by meiosis-activated protein kinase p44(mpk).
A; Reference number: S12904; MUID: 91032186; PMID: 1699809
A; Accession: S12904
A; Status: preliminary
A; Molecule type: protein
A; Residues: 1-14 <SAN>
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gamma 1 gliadin - wheat (fragment)
C; Species: Triticum sp. (wheat)
C;Date: 08-Dec-1995 #sequence revision 08-Dec-1995 #text change 08-Dec-1995
C; Accession: A56891
R; Sjostrom, H.; Friis, S.U.; Noren, O.; Anthonsen, D.
Clin. Chim. Acta 207, 227-237, 1992
A; Title: Purification and characterisation of antigenic gliadins in coeliac
A; Reference number: A56891; MUID: 93009000; PMID: 1395028
A; Contents: Kadett
A; Accession: A56891
A; Status: preliminary
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N; Alternate names: superprotein peptide 9
C; Species: Oryctolagus cuniculus (domestic rabbit)
C;Date: 31-Mar-1993 #sequence_revision 18-Nov-1994 #text_change 15-Jun-1996
C; Accession: F44823
R; Loewy, A.; Liu, W.S.; Baitinger, C.; Willard, M.B.
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J. Neurosci. 11, 3412-3421, 1991
A; Title: The major 35S-methionine-labeled rapidly transported protein
(superprotein) is identical to SNAP-25, a protein of synaptic terminals.
A; Reference number: A44823; MUID: 92044785; PMID: 1941090
A; Accession: F44823
A; Status: preliminary
A; Molecule type: protein
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crystal protein, 28K - Bacillus thuringiensis (fragment)
C; Species: Bacillus thuringiensis
C; Date: 22-Jul-1987 #sequence revision 22-Jul-1987 #text change 20-Jun-2000
C; Accession: A24099
R; Chestukhina, G.G.; Zalunin, I.A.; Kostina, L.I.; Bormatova, M.E.; Klepikova,
F.S.; Khodova, O.M.; Stepanov, V.M.
FEBS Lett. 190, 345-348, 1985
A; Title: Structural features of crystal-forming proteins produced by Bacillus
thuringiensis subspecies israelensis.
A; Reference number: A91347
A; Accession: A24099
A; Molecule type: protein
A; Residues: 1-16 < CHE>
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PH0763
T-cell receptor beta chain (F15) - mouse (fragment)
C; Species: Mus musculus (house mouse)
C;Date: 17-Jul-1992 #sequence revision 17-Jul-1992 #text change 05-Nov-1999
C; Accession: PH0763
R; Casanova, J.L.; Romero, P.; Widmann, C.; Kourilsky, P.; Maryanski, J.L.
J. Exp. Med. 174, 1371-1383, 1991
A; Title: T cell receptor genes in a series of class I major histocompatibility
complex-restricted cytotoxic T lymphocyte clones specific for a Plasmodium
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berghei nonapeptide: implications for T cell allelic exclusion and antigen-

specific repertoire.

A; Reference number: PH0746; MUID: 92078846; PMID: 1836010

A; Accession: PH0763 A; Molecule type: mRNA A; Residues: 1-16 <CAS>

A; Cross-references: EMBL: X60857; NID: g50933; PIDN: CAA43247.1; PID: g50934

A; Experimental source: T lymphocyte

C; Keywords: T-cell receptor

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Search completed: July 4, 2004, 04:47:14

Job time : 15.0672 secs

## GenCore version 5.1.6 Copyright (c) 1993 - 2004 Compugen Ltd.

OM protein - protein search, using sw model

Run on: July 4, 2004, 04:45:52; Search time 33.3657 Seconds

(without alignments)

158.601 Million cell updates/sec

Title: US-09-641-802-2

Perfect score: 89

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Gapop 10.0 , Gapext 0.5

Searched: 1276540 seqs, 311283816 residues

Total number of hits satisfying chosen parameters: 203405

Minimum DB seq length: 7
Maximum DB seq length: 18

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 100 summaries

Database: Published Applications AA:\*

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Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

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Result Query

No. Score Match Length DB ID

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4	30	33.7	16	12		Sequence 130, App
5	29	32.6	9	13	US-10-017-327-3	Sequence 3, Appli
6	29	32.6	10	14	US-10-073-054-20	Sequence 20, Appl
7	29	32.6	13	12	US-10-469-101-53	Sequence 53, Appl
8	29	32.6	18	9	US-09-864-761-40147	Sequence 40147, A
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63	26	29.2	15	16	US-10-409-613-121	Sequence 121, App
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72	25	28.1	10	12	US-10-325-810-116	Sequence 116, App
73	25	28.1	12	L 4	US-10-193-477-146	Sequence 146, App
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## ALIGNMENTS

## RESULT 1 US-10-281-652-2

- ; Sequence 2, Application US/10281652
- ; Publication No. US20030091606A1
- ; GENERAL INFORMATION:
- ; APPLICANT: STANTON, G. John
- ; APPLICANT: HUGHES, Thomas K. ; APPLICANT: BOLDOGH, Istvan

```
TITLE OF INVENTION: USE OF COLOSTRININ, CONSTITUENT PEPTIDES THEREOF, AND
  TITLE OF INVENTION: ANALOGS THEREOF AS OXIDATIVE STRESS REGULATORS
; FILE REFERENCE: 265.00220101
  CURRENT APPLICATION NUMBER: US/10/281,652
  CURRENT FILING DATE: 2002-10-28
  PRIOR APPLICATION NUMBER: US/09/641,803
 PRIOR FILING DATE: 2000-08-17
; PRIOR APPLICATION NUMBER: 60/149,310
 PRIOR FILING DATE: 1999-08-17
; NUMBER OF SEQ ID NOS: 34
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; SEQ ID NO 2
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   TYPE: PRT
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   OTHER INFORMATION: Description of Artificial Sequence: synthetic
   OTHER INFORMATION: peptide
US-10-281-652-2
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US-10-286-457-195
; Sequence 195, Application US/10286457
; Publication No. US20030166004A1
; GENERAL INFORMATION:
  APPLICANT: JENO GYURIS et al.
  TITLE OF INVENTION: ENDOTHELIAL-CELL BINDING PEPTIDES FOR DIAGNOSIS AND
THERAPY
; FILE REFERENCE: GPCI-P01-178
  CURRENT APPLICATION NUMBER: US/10/286,457
; CURRENT FILING DATE: 2002-11-01
; PRIOR APPLICATION NUMBER: 60/334822
 PRIOR FILING DATE: 2001-11-01
; NUMBER OF SEQ ID NOS: 684
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; Publication No. US20030091606A1
; GENERAL INFORMATION:
  APPLICANT: STANTON, G. John
  APPLICANT: HUGHES, Thomas K.
  APPLICANT: BOLDOGH, Istvan
  TITLE OF INVENTION: USE OF COLOSTRININ, CONSTITUENT PEPTIDES THEREOF, AND
  TITLE OF INVENTION: ANALOGS THEREOF AS OXIDATIVE STRESS REGULATORS
  FILE REFERENCE: 265.00220101
  CURRENT APPLICATION NUMBER: US/10/281,652
  CURRENT FILING DATE: 2002-10-28
  PRIOR APPLICATION NUMBER: US/09/641,803
  PRIOR FILING DATE: 2000-08-17
  PRIOR APPLICATION NUMBER: 60/149,310
  PRIOR FILING DATE: 1999-08-17
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; Sequence 130, Application US/10346162
; Publication No. US20030224390A1
; GENERAL INFORMATION:
  APPLICANT: KARO BIO USA, INC.
  APPLICANT: FOWLKES, Dana M.
  APPLICANT: BARNETT, Thomas R.
  APPLICANT: BUEHRER, Benjamin
  TITLE OF INVENTION: METHOD OF IDENTIFYING CONFORMATION-SENSITIVE BINDING
PEPTIDES AND USES
; TITLE OF INVENTION: THEREOF
; FILE REFERENCE: PAIGE=1H
; CURRENT APPLICATION NUMBER: US/10/346,162
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  PRIOR APPLICATION NUMBER: US 09/614,865
  PRIOR FILING DATE: 2000-07-12
; PRIOR APPLICATION NUMBER: US 09/860,688
  PRIOR FILING DATE: 2001-05-21
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; Publication No. US20020155471A1
; GENERAL INFORMATION:
; APPLICANT: Charles A. Nicolette
; TITLE OF INVENTION: COMPOUNDS FOR THERAPY AND DIAGNOSIS AND
; TITLE OF INVENTION: METHODS FOR USING SAME
; FILE REFERENCE: GZ 2101.20
  CURRENT APPLICATION NUMBER: US/10/017,327
  CURRENT FILING DATE: 2001-12-06
; NUMBER OF SEQ ID NOS: 11
; SOFTWARE: FastSEQ for Windows Version 4.0
; SEQ ID NO 3
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   TYPE: PRT
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Qу
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RESULT 6
US-10-073-054-20
; Sequence 20, Application US/10073054
; Publication No. US20030167485A1
; GENERAL INFORMATION:
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; APPLICANT: Garvan Institute of Medical Research
; APPLICANT: HERZOG, Herbert
; TITLE OF INVENTION: No. US20030167485Alel G-protein coupled receptor-encoding
gene and diagnostic uses
; TITLE OF INVENTION: therefor
  FILE REFERENCE: 1871-132 (93702-CIP/MRO)
  CURRENT APPLICATION NUMBER: US/10/073,054
; CURRENT FILING DATE: 2002-02-12
; PRIOR APPLICATION NUMBER: US 09/308,696
 PRIOR FILING DATE: 1999-06-11
 PRIOR APPLICATION NUMBER: PCT/AU98/00805
 PRIOR FILING DATE: 1998-09-24
; PRIOR APPLICATION NUMBER: AU PO9386
; PRIOR FILING DATE: 1997-09-24
; NUMBER OF SEQ ID NOS: 20
; SOFTWARE: PatentIn version 3.1
; SEQ ID NO 20
   LENGTH: 10
   TYPE: PRT
   ORGANISM: N-terminal peptide
US-10-073-054-20
                         32.6%; Score 29; DB 14; Length 10;
 Query Match
 Best Local Similarity 85.7%; Pred. No. 6.1e+02;
           6; Conservative 0; Mismatches 1; Indels 0; Gaps
 Matches
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          . 3 TPQPLLQ 9
Qу
             111 111
           2 TPQSLLQ 8
Db
RESULT 7
US-10-469-101-53
; Sequence 53, Application US/10469101
; Publication No. US20040071688A1
; GENERAL INFORMATION:
; APPLICANT: CARR, Francis J.
; APPLICANT: CARTER, Graham
  TITLE OF INVENTION: MODIFIED THROMBOPOIETIN WITH REDUCED
  TITLE OF INVENTION: IMMUNOGENICITY
 FILE REFERENCE: MER-109
  CURRENT APPLICATION NUMBER: US/10/469,101
  CURRENT FILING DATE: 2002-08-26
; PRIOR APPLICATION NUMBER: EP 01104702.4
; PRIOR FILING DATE: 2001-02-26
  PRIOR APPLICATION NUMBER: PCT/EP02/01931
  PRIOR FILING DATE: 2002-02-22
 NUMBER OF SEQ ID NOS: 126
  SOFTWARE: FastSEQ for Windows Version 4.0
; SEQ ID NO 53
   LENGTH: 13
   TYPE: PRT
   ORGANISM: Artificial Sequence
   OTHER INFORMATION: MHC class II binding epitope
US-10-469-101-53
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Query Match
                          32.6%; Score 29; DB 12; Length 13;
  Best Local Similarity 50.0%; Pred. No. 8.1e+02;
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            6; Conservative
                                                 5; Indels
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            5 QPLLQVMMEPQG 16
Qу
              + +
                    : |||
Db
            1 QSLLGTQLPPQG 12
RESULT 8
US-09-864-761-40147
; Sequence 40147, Application US/09864761
; Patent No. US20020048763A1
; GENERAL INFORMATION:
; APPLICANT: Penn, Sharron G.
  APPLICANT: Rank, David R.
  APPLICANT: Hanzel, David K.
  APPLICANT: Chen, Wensheng
  TITLE OF INVENTION: HUMAN GENOME-DERIVED SINGLE EXON NUCLEIC ACID PROBES
USEFUL FOR
  TITLE OF INVENTION: GENE EXPRESSION ANALYSIS BY MICROARRAY
   FILE REFERENCE: Aeomica-X-1
   CURRENT APPLICATION NUMBER: US/09/864,761
   CURRENT FILING DATE: 2001-05-23
   PRIOR APPLICATION NUMBER: US 60/180,312
   PRIOR FILING DATE: 2000-02-04
   PRIOR APPLICATION NUMBER: US 60/207,456
   PRIOR FILING DATE: 2000-05-26
   PRIOR APPLICATION NUMBER: US 09/632,366
   PRIOR FILING DATE: 2000-08-03
   PRIOR APPLICATION NUMBER: GB 24263.6
   PRIOR FILING DATE: 2000-10-04
   PRIOR APPLICATION NUMBER: US 60/236,359
   PRIOR FILING DATE: 2000-09-27
   PRIOR APPLICATION NUMBER: PCT/US01/00666
   PRIOR FILING DATE: 2001-01-30
   PRIOR APPLICATION NUMBER: PCT/US01/00667
   PRIOR FILING DATE: 2001-01-30
   PRIOR APPLICATION NUMBER: PCT/US01/00664
   PRIOR FILING DATE: 2001-01-30
   PRIOR APPLICATION NUMBER: PCT/US01/00669
   PRIOR FILING DATE: 2001-01-30
   PRIOR APPLICATION NUMBER: PCT/US01/00665
   PRIOR FILING DATE: 2001-01-30
   PRIOR APPLICATION NUMBER: PCT/US01/00668
   PRIOR FILING DATE: 2001-01-30
   PRIOR APPLICATION NUMBER: PCT/US01/00663
   PRIOR FILING DATE: 2001-01-30
   PRIOR APPLICATION NUMBER: PCT/US01/00662
   PRIOR FILING DATE: 2001-01-30
   PRIOR APPLICATION NUMBER: PCT/US01/00661
   PRIOR FILING DATE: 2001-01-30
   PRIOR APPLICATION NUMBER: PCT/US01/00670
   PRIOR FILING DATE: 2001-01-30
   PRIOR APPLICATION NUMBER: US 60/234,687
   PRIOR FILING DATE: 2000-09-21
   PRIOR APPLICATION NUMBER: US 09/608,408
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PRIOR FILING DATE: 2000-06-30
  PRIOR APPLICATION NUMBER: US 09/774,203
  PRIOR FILING DATE: 2001-01-29
  NUMBER OF SEQ ID NOS: 49117
  SOFTWARE: Annomax Sequence Listing Engine vers. 1.1
 SEQ ID NO 40147
   LENGTH: 18
   TYPE: PRT
   ORGANISM: Homo sapiens
   FEATURE:
   OTHER INFORMATION: MAP TO AC005343.1
   OTHER INFORMATION: EXPRESSED IN HELA, SIGNAL = 3.8
   OTHER INFORMATION: EXPRESSED IN FETAL LIVER, SIGNAL =
   OTHER INFORMATION: EXPRESSED IN BRAIN, SIGNAL = 4.3
   OTHER INFORMATION: EXPRESSED IN BONE MARROW, SIGNAL =
   OTHER INFORMATION: EXPRESSED IN ADULT LIVER, SIGNAL = 4.8
   OTHER INFORMATION: EXPRESSED IN PLACENTA, SIGNAL = 3.8
   OTHER INFORMATION: EXPRESSED IN HEART, SIGNAL = 3.3
    OTHER INFORMATION: EXPRESSED IN LUNG, SIGNAL = 4.2
US-09-864-761-40147
                         32.6%; Score 29; DB 9; Length 18;
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  Best Local Similarity 100.0%; Pred. No. 1.1e+03;
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           6; Conservative
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                                                0; Indels
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           5 QPLLQV 10
Qу
             11111
Db
           3 OPLLOV 8
RESULT 9
US-10-013-312-33
; Sequence 33, Application US/10013312
; Publication No. US20030223990A1
; GENERAL INFORMATION:
; APPLICANT: RAITANO, ARTHUR
; APPLICANT: CHALLITA-EID, PIA
; APPLICANT: FARIS, MARY
; APPLICANT: HUBERT, RENE
  APPLICANT: GE, WANGMAO
  APPLICANT: JAKOBOVITZ, AYA
  TITLE OF INVENTION: NUCLEIC ACID AND CORRESPONDING PROTEIN ENTITLED
  TITLE OF INVENTION: 193P1E1B USEFUL IN TREATMENT AND DETECTION OF CANCER
  FILE REFERENCE: 51158-20063.00
; CURRENT APPLICATION NUMBER: US/10/013,312
; CURRENT FILING DATE: 2002-05-30
; NUMBER OF SEQ ID NOS: 3005
  SOFTWARE: PatentIn Ver. 2.1
; SEQ ID NO 33
   LENGTH: 9
   TYPE: PRT
   ORGANISM: Homo sapiens
US-10-013-312-33
 Query Match
                         31.5%; Score 28; DB 12; Length 9;
 Best Local Similarity 57.1%; Pred. No. 1.2e+06;
 Matches 4; Conservative 2; Mismatches 1; Indels 0; Gaps
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3 TPQPLLQ 9
Qy
             11 1::1
           3 TPSPIIQ 9
Db
RESULT 10
US-10-013-312-412
; Sequence 412, Application US/10013312
; Publication No. US20030223990A1
; GENERAL INFORMATION:
; APPLICANT: RAITANO, ARTHUR
; APPLICANT: CHALLITA-EID, PIA
; APPLICANT: FARIS, MARY
; APPLICANT: HUBERT, RENE
; APPLICANT: GE, WANGMAO
 APPLICANT: JAKOBOVITZ, AYA
  TITLE OF INVENTION: NUCLEIC ACID AND CORRESPONDING PROTEIN ENTITLED
  TITLE OF INVENTION: 193P1E1B USEFUL IN TREATMENT AND DETECTION OF CANCER
; FILE REFERENCE: 51158-20063.00
; CURRENT APPLICATION NUMBER: US/10/013,312
  CURRENT FILING DATE: 2002-05-30
 NUMBER OF SEQ ID NOS: 3005
 SOFTWARE: PatentIn Ver. 2.1
; SEQ ID NO 412
   LENGTH: 9
   TYPE: PRT
   ORGANISM: Homo sapiens
US-10-013-312-412
                         31.5%; Score 28; DB 12; Length 9;
  Query Match
  Best Local Similarity 57.1%; Pred. No. 1.2e+06;
          4; Conservative 2; Mismatches 1; Indels 0; Gaps
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 Matches
Qy
           3 TPOPLLO 9
             || |::|
           1 TPSPIIQ 7
Db
RESULT 11
US-10-013-312-502
; Sequence 502, Application US/10013312
; Publication No. US20030223990A1
; GENERAL INFORMATION:
; APPLICANT: RAITANO, ARTHUR
; APPLICANT: CHALLITA-EID, PIA
; APPLICANT: FARIS, MARY
; APPLICANT: HUBERT, RENE
  APPLICANT: GE, WANGMAO
  APPLICANT: JAKOBOVITZ, AYA
  TITLE OF INVENTION: NUCLEIC ACID AND CORRESPONDING PROTEIN ENTITLED
  TITLE OF INVENTION: 193P1E1B USEFUL IN TREATMENT AND DETECTION OF CANCER
; FILE REFERENCE: 51158-20063.00
; CURRENT APPLICATION NUMBER: US/10/013,312
; CURRENT FILING DATE: 2002-05-30
; NUMBER OF SEQ ID NOS: 3005
; SOFTWARE: PatentIn Ver. 2.1
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; SEQ ID NO 502
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US-10-013-312-502
                         31.5%; Score 28; DB 12; Length 9;
  Query Match
  Best Local Similarity 57.1%; Pred. No. 1.2e+06;
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                               2; Mismatches
                                                                           0;
           4; Conservative
                                               1; Indels
           3 TPQPLLQ 9
Qу
             11 1::1
Db
           1 TPSPIIO 7
RESULT 12
US-10-013-312-605
; Sequence 605, Application US/10013312
; Publication No. US20030223990A1
; GENERAL INFORMATION:
  APPLICANT: RAITANO, ARTHUR
  APPLICANT: CHALLITA-EID, PIA
  APPLICANT: FARIS, MARY
  APPLICANT: HUBERT, RENE
  APPLICANT: GE, WANGMAO
  APPLICANT: JAKOBOVITZ, AYA
  TITLE OF INVENTION: NUCLEIC ACID AND CORRESPONDING PROTEIN ENTITLED
  TITLE OF INVENTION: 193P1E1B USEFUL IN TREATMENT AND DETECTION OF CANCER
  FILE REFERENCE: 51158-20063.00
  CURRENT APPLICATION NUMBER: US/10/013,312
  CURRENT FILING DATE: 2002-05-30
; NUMBER OF SEQ ID NOS: 3005
  SOFTWARE: PatentIn Ver. 2.1
; SEQ ID NO 605
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    ORGANISM: Homo sapiens
US-10-013-312-605
  Query Match
                         31.5%; Score 28; DB 12; Length 9;
                         57.1%; Pred. No. 1.2e+06;
  Best Local Similarity
                               2; Mismatches
                                                1; Indels
                                                               0; Gaps
                                                                           0;
  Matches
            4; Conservative
            3 TPQPLLQ 9
Qу
             11 1::1
            1 TPSPIIQ 7
Db
RESULT 13
US-10-013-312-1358
; Sequence 1358, Application US/10013312
; Publication No. US20030223990A1
; GENERAL INFORMATION:
; APPLICANT: RAITANO, ARTHUR
; APPLICANT: CHALLITA-EID, PIA
; APPLICANT: FARIS, MARY
; APPLICANT: HUBERT, RENE
```

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; APPLICANT: GE, WANGMAO
; APPLICANT: JAKOBOVITZ, AYA
  TITLE OF INVENTION: NUCLEIC ACID AND CORRESPONDING PROTEIN ENTITLED
  TITLE OF INVENTION: 193P1E1B USEFUL IN TREATMENT AND DETECTION OF CANCER
  FILE REFERENCE: 51158-20063.00
  CURRENT APPLICATION NUMBER: US/10/013,312
  CURRENT FILING DATE: 2002-05-30
; NUMBER OF SEQ ID NOS: 3005
; SOFTWARE: PatentIn Ver. 2.1
; SEQ ID NO 1358
   LENGTH: 9
   TYPE: PRT
   ORGANISM: Homo sapiens
US-10-013-312-1358
 Query Match
                         31.5%; Score 28; DB 12; Length 9;
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           4; Conservative 2; Mismatches 1; Indels 0; Gaps
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 Matches
           3 TPQPLLQ 9
Qу
             || |::|
           1 TPSPIIQ 7
Db
RESULT 14
US-10-013-312-1522
; Sequence 1522, Application US/10013312
; Publication No. US20030223990A1
; GENERAL INFORMATION:
; APPLICANT: RAITANO, ARTHUR
; APPLICANT: CHALLITA-EID, PIA
; APPLICANT: FARIS, MARY
; APPLICANT: HUBERT, RENE
  APPLICANT: GE, WANGMAO
  APPLICANT: JAKOBOVITZ, AYA
  TITLE OF INVENTION: NUCLEIC ACID AND CORRESPONDING PROTEIN ENTITLED
  TITLE OF INVENTION: 193P1E1B USEFUL IN TREATMENT AND DETECTION OF CANCER
  FILE REFERENCE: 51158-20063.00
  CURRENT APPLICATION NUMBER: US/10/013,312
  CURRENT FILING DATE: 2002-05-30
; NUMBER OF SEQ ID NOS: 3005
  SOFTWARE: PatentIn Ver. 2.1
; SEQ ID NO 1522
   LENGTH: 9
   TYPE: PRT
   ORGANISM: Homo sapiens
US-10-013-312-1522
                         31.5%; Score 28; DB 12; Length 9;
  Query Match
  Best Local Similarity 57.1%; Pred. No. 1.2e+06;
          4; Conservative 2; Mismatches 1; Indels 0; Gaps
  Matches
Qу
           3 TPQPLLQ 9
             11 1::1
Db
           2 TPSPIIQ 8
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RESULT 15
US-10-013-312-1598
; Sequence 1598, Application US/10013312
; Publication No. US20030223990A1
; GENERAL INFORMATION:
 APPLICANT: RAITANO, ARTHUR
; APPLICANT: CHALLITA-EID, PIA
; APPLICANT: FARIS, MARY
; APPLICANT: HUBERT, RENE
; APPLICANT: GE, WANGMAO
 APPLICANT: JAKOBOVITZ, AYA
  TITLE OF INVENTION: NUCLEIC ACID AND CORRESPONDING PROTEIN ENTITLED
  TITLE OF INVENTION: 193P1E1B USEFUL IN TREATMENT AND DETECTION OF CANCER
; FILE REFERENCE: 51158-20063.00
; CURRENT APPLICATION NUMBER: US/10/013,312
; CURRENT FILING DATE: 2002-05-30
; NUMBER OF SEQ ID NOS: 3005
; SOFTWARE: PatentIn Ver. 2.1
; SEQ ID NO 1598
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   TYPE: PRT
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US-10-013-312-1598
                         31.5%; Score 28; DB 12; Length 9;
  Query Match
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  Matches 4; Conservative 2; Mismatches 1; Indels 0; Gaps
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           3 TPQPLLQ 9
Qу
             1 TPSPIIQ 7
Db
RESULT 16
US-10-013-312-1693
; Sequence 1693, Application US/10013312
; Publication No. US20030223990A1
; GENERAL INFORMATION:
; APPLICANT: RAITANO, ARTHUR
; APPLICANT: CHALLITA-EID, PIA
 APPLICANT: FARIS, MARY
  APPLICANT: HUBERT, RENE
  APPLICANT: GE, WANGMAO
; APPLICANT: JAKOBOVITZ, AYA
  TITLE OF INVENTION: NUCLEIC ACID AND CORRESPONDING PROTEIN ENTITLED
  TITLE OF INVENTION: 193P1E1B USEFUL IN TREATMENT AND DETECTION OF CANCER
; FILE REFERENCE: 51158-20063.00
; CURRENT APPLICATION NUMBER: US/10/013,312
  CURRENT FILING DATE: 2002-05-30
  NUMBER OF SEQ ID NOS: 3005
  SOFTWARE: PatentIn Ver. 2.1
; SEQ ID NO 1693
   LENGTH: 9
   TYPE: PRT
   ORGANISM: Homo sapiens
US-10-013-312-1693
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Query Match
                         31.5%; Score 28; DB 12; Length 9;
  Best Local Similarity 57.1%; Pred. No. 1.2e+06;
           4; Conservative 2; Mismatches
                                                1; Indels 0; Gaps
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           3 TPQPLLQ 9
Qу
              11 1::1
Db
           1 TPSPIIQ 7
RESULT 17
US-10-013-312-1721
; Sequence 1721, Application US/10013312
; Publication No. US20030223990A1
; GENERAL INFORMATION:
; APPLICANT: RAITANO, ARTHUR
; APPLICANT: CHALLITA-EID, PIA
; APPLICANT: FARIS, MARY
  APPLICANT: HUBERT, RENE
  APPLICANT: GE, WANGMAO
  APPLICANT: JAKOBOVITZ, AYA
  TITLE OF INVENTION: NUCLEIC ACID AND CORRESPONDING PROTEIN ENTITLED
  TITLE OF INVENTION: 193P1E1B USEFUL IN TREATMENT AND DETECTION OF CANCER
  FILE REFERENCE: 51158-20063.00
  CURRENT APPLICATION NUMBER: US/10/013,312
  CURRENT FILING DATE: 2002-05-30
; NUMBER OF SEQ ID NOS: 3005
  SOFTWARE: PatentIn Ver. 2.1
; SEQ ID NO 1721
   LENGTH: 9
   TYPE: PRT
   ORGANISM: Homo sapiens
US-10-013-312-1721
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            3 TPQPLLQ 9
Qу
             11 1::1
Db
           1 TPSPIIQ 7
RESULT 18
US-10-013-312-1810
; Sequence 1810, Application US/10013312
; Publication No. US20030223990A1
; GENERAL INFORMATION:
; APPLICANT: RAITANO, ARTHUR
  APPLICANT: CHALLITA-EID, PIA
  APPLICANT: FARIS, MARY
  APPLICANT: HUBERT, RENE
; APPLICANT: GE, WANGMAO
  APPLICANT: JAKOBOVITZ, AYA
  TITLE OF INVENTION: NUCLEIC ACID AND CORRESPONDING PROTEIN ENTITLED
; TITLE OF INVENTION: 193P1E1B USEFUL IN TREATMENT AND DETECTION OF CANCER
; FILE REFERENCE: 51158-20063.00
; CURRENT APPLICATION NUMBER: US/10/013,312
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; CURRENT FILING DATE: 2002-05-30
; NUMBER OF SEQ ID NOS: 3005
; SOFTWARE: PatentIn Ver. 2.1
; SEQ ID NO 1810
   LENGTH: 9
   TYPE: PRT
   ORGANISM: Homo sapiens
US-10-013-312-1810
                         31.5%; Score 28; DB 12; Length 9;
 Query Match
                         57.1%; Pred. No. 1.2e+06;
 Best Local Similarity
                               2; Mismatches 1; Indels
 Matches
           4; Conservative
                                                             0; Gaps
           3 TPQPLLQ 9
Qу
             11 1::1
           1 TPSPIIO 7
Db
RESULT 19
US-10-013-312-1928
; Sequence 1928, Application US/10013312
; Publication No. US20030223990A1
; GENERAL INFORMATION:
; APPLICANT: RAITANO, ARTHUR
 APPLICANT: CHALLITA-EID, PIA
  APPLICANT: FARIS, MARY
  APPLICANT: HUBERT, RENE
; APPLICANT: GE, WANGMAO
; APPLICANT: JAKOBOVITZ, AYA
  TITLE OF INVENTION: NUCLEIC ACID AND CORRESPONDING PROTEIN ENTITLED
  TITLE OF INVENTION: 193P1E1B USEFUL IN TREATMENT AND DETECTION OF CANCER
; FILE REFERENCE: 51158-20063.00
; CURRENT APPLICATION NUMBER: US/10/013,312
  CURRENT FILING DATE: 2002-05-30
  NUMBER OF SEQ ID NOS: 3005
  SOFTWARE: PatentIn Ver. 2.1
; SEQ ID NO 1928
   LENGTH: 9
   TYPE: PRT
   ORGANISM: Homo sapiens
US-10-013-312-1928
  Query Match
                         31.5%; Score 28; DB 12; Length 9;
  Best Local Similarity 57.1%; Pred. No. 1.2e+06;
 Matches 4; Conservative 2; Mismatches 1; Indels
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                                                                           0;
           3 TPQPLLQ 9
Qу
             11 1::1
           1 TPSPIIQ 7
Db
RESULT 20
US-10-013-312-1983
; Sequence 1983, Application US/10013312
; Publication No. US20030223990A1
; GENERAL INFORMATION:
; APPLICANT: RAITANO, ARTHUR
```

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; APPLICANT: CHALLITA-EID, PIA
; APPLICANT: FARIS, MARY
; APPLICANT: HUBERT, RENE
; APPLICANT: GE, WANGMAO
  APPLICANT: JAKOBOVITZ, AYA
  TITLE OF INVENTION: NUCLEIC ACID AND CORRESPONDING PROTEIN ENTITLED
  TITLE OF INVENTION: 193P1E1B USEFUL IN TREATMENT AND DETECTION OF CANCER
; FILE REFERENCE: 51158-20063.00
  CURRENT APPLICATION NUMBER: US/10/013,312
; CURRENT FILING DATE: 2002-05-30
; NUMBER OF SEQ ID NOS: 3005
  SOFTWARE: PatentIn Ver. 2.1
; SEQ ID NO 1983
  LENGTH: 9
   TYPE: PRT
   ORGANISM: Homo sapiens
US-10-013-312-1983
 Query Match 31.5%; Score 28; DB 12; Length 9; Best Local Similarity 57.1%; Pred. No. 1.2e+06;
 Matches 4; Conservative 2; Mismatches 1; Indels 0; Gaps
                                                                            0;
           3 TPQPLLQ 9
Qу
             | | | | ::|
Db
           1 TPSPIIQ 7
RESULT 21
US-10-013-312-2020
; Sequence 2020, Application US/10013312
; Publication No. US20030223990A1
; GENERAL INFORMATION:
; APPLICANT: RAITANO, ARTHUR
; APPLICANT: CHALLITA-EID, PIA
  APPLICANT: FARIS, MARY
; APPLICANT: HUBERT, RENE
; APPLICANT: GE, WANGMAO
; APPLICANT: JAKOBOVITZ, AYA
  TITLE OF INVENTION: NUCLEIC ACID AND CORRESPONDING PROTEIN ENTITLED
  TITLE OF INVENTION: 193P1E1B USEFUL IN TREATMENT AND DETECTION OF CANCER
 FILE REFERENCE: 51158-20063.00
  CURRENT APPLICATION NUMBER: US/10/013,312
  CURRENT FILING DATE: 2002-05-30
; NUMBER OF SEQ ID NOS: 3005
; SOFTWARE: PatentIn Ver. 2.1
; SEQ ID NO 2020
   LENGTH: 9
   TYPE: PRT
   ORGANISM: Homo sapiens
US-10-013-312-2020
  Query Match
                         31.5%; Score 28; DB 12; Length 9;
  Best Local Similarity 57.1%; Pred. No. 1.2e+06;
 Matches 4; Conservative 2; Mismatches 1; Indels 0; Gaps
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Qу
           3 TPQPLLQ 9
             11 1::1
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RESULT 22
US-09-908-322-53
; Sequence 53, Application US/09908322
; Patent No. US20020107194A1
    GENERAL INFORMATION:
         APPLICANT: Ish-Horowicz, David
                    Henrique, Domingos Manuel Pinto
                    Lewis, Julian Hart
                    Artavanis-Tsakonas, Spyridon
                    Gray, Grace
         TITLE OF INVENTION: NUCLEOTIDE AND PROTEIN SEQUENCES OF
                             VERTEBRATE DELTA GENE AND METHODS BASED THEREON
         NUMBER OF SEQUENCES: 94
         CORRESPONDENCE ADDRESS:
              ADDRESSEE: Pennie & Edmonds LLP
              STREET: 1155 Avenue of the Americas
              CITY: New York
              STATE: NY
              COUNTRY: USA
              ZIP: 10036/2711
         COMPUTER READABLE FORM:
              MEDIUM TYPE: Diskette
              COMPUTER: IBM Compatible
              OPERATING SYSTEM: DOS
              SOFTWARE: FastSEQ Version 2.0
         CURRENT APPLICATION DATA:
              APPLICATION NUMBER: US/09/908,322
              FILING DATE: 17-Jul-2001
              CLASSIFICATION: <Unknown>
         PRIOR APPLICATION DATA:
              APPLICATION NUMBER: 08/981,392
              FILING DATE: 22-DEC-1997
         ATTORNEY/AGENT INFORMATION:
              NAME: Misrock, S Leslie
              REGISTRATION NUMBER: 18,872
              REFERENCE/DOCKET NUMBER: 7326-123
         TELECOMMUNICATION INFORMATION:
              TELEPHONE: 212-790-9090
              TELEFAX: 212-869-8864
              TELEX: 66141 PENNIE
    INFORMATION FOR SEQ ID NO: 53:
         SEQUENCE CHARACTERISTICS:
              LENGTH: 10 amino acids
              TYPE: amino acid
              STRANDEDNESS: <Unknown>
              TOPOLOGY: unknown
         MOLECULE TYPE: peptide
         SEQUENCE DESCRIPTION: SEQ ID NO: 53:
US-09-908-322-53
                          31.5%; Score 28; DB 9; Length 10;
  Query Match
  Best Local Similarity 50.0%; Pred. No. 8.7e+02;
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            5; Conservative 2; Mismatches 3; Indels
  Matches
                                                                             0;
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4 PQPLLQVMME 13
Qу
             1111::
Db
            1 PQPLVRTEQE 10
RESULT 23
US-09-783-931-53
; Sequence 53, Application US/09783931
; Publication No. US20030073620A1
   GENERAL INFORMATION:
        APPLICANT: Ish-Horowicz, David
                    Henrique, Domingos Manuel Pinto
                    Lewis, Julian Hart
                    Artavanis-Tsakonas, Spyridon
                    Gray, Grace
         TITLE OF INVENTION: ANTIBODIES TO VERTEBRATE DELTA PROTEINS
                             AND FRAGMENTS
        NUMBER OF SEQUENCES: 94
         CORRESPONDENCE ADDRESS:
              ADDRESSEE: Pennie & Edmonds LLP
              STREET: 1155 Avenue of the Americas
              CITY: New York
              STATE: NY
              COUNTRY: USA
              ZIP: 10036/2711
        COMPUTER READABLE FORM:
              MEDIUM TYPE: Diskette
              COMPUTER: IBM Compatible
              OPERATING SYSTEM: DOS
              SOFTWARE: FastSEQ Version 2.0
         CURRENT APPLICATION DATA:
              APPLICATION NUMBER: US/09/783,931
              FILING DATE: 15-Feb-2001
              CLASSIFICATION: <Unknown>
         PRIOR APPLICATION DATA:
              APPLICATION NUMBER: 08/981,392
              FILING DATE: 22-DEC-1997
         ATTORNEY/AGENT INFORMATION:
              NAME: Antler, Adriane M.
              REGISTRATION NUMBER: 32,605
              REFERENCE/DOCKET NUMBER: 7326-122
         TELECOMMUNICATION INFORMATION:
              TELEPHONE: 212-790-9090
              TELEFAX: 212-869-8864
              TELEX: 66141 PENNIE
    INFORMATION FOR SEQ ID NO: 53:
         SEQUENCE CHARACTERISTICS:
              LENGTH: 10 amino acids
              TYPE: amino acid
              STRANDEDNESS: <Unknown>
              TOPOLOGY: unknown
         MOLECULE TYPE: peptide
         SEQUENCE DESCRIPTION: SEQ ID NO: 53:
US-09-783-931-53
  Query Match
                          31.5%; Score 28; DB 10; Length 10;
  Best Local Similarity 50.0%; Pred. No. 8.7e+02;
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Matches
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Qу
           4 PQPLLQVMME 13
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           1 PQPLVRTEQE 10
RESULT 24
US-10-013-312-82
; Sequence 82, Application US/10013312
; Publication No. US20030223990A1
; GENERAL INFORMATION:
; APPLICANT: RAITANO, ARTHUR
; APPLICANT: CHALLITA-EID, PIA
; APPLICANT: FARIS, MARY
; APPLICANT: HUBERT, RENE
; APPLICANT: GE, WANGMAO
 APPLICANT: JAKOBOVITZ, AYA
  TITLE OF INVENTION: NUCLEIC ACID AND CORRESPONDING PROTEIN ENTITLED
  TITLE OF INVENTION: 193P1E1B USEFUL IN TREATMENT AND DETECTION OF CANCER
 FILE REFERENCE: 51158-20063.00
; CURRENT APPLICATION NUMBER: US/10/013,312
; CURRENT FILING DATE: 2002-05-30
; NUMBER OF SEQ ID NOS: 3005
 SOFTWARE: PatentIn Ver. 2.1
; SEQ ID NO 82
   LENGTH: 10
   TYPE: PRT
   ORGANISM: Homo sapiens
US-10-013-312-82
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                         31.5%; Score 28; DB 12; Length 10;
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          4; Conservative 2; Mismatches 1; Indels
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Db
           3 TPSPIIQ 9
RESULT 25
US-10-013-312-196
; Sequence 196, Application US/10013312
; Publication No. US20030223990A1
; GENERAL INFORMATION:
; APPLICANT: RAITANO, ARTHUR
; APPLICANT: CHALLITA-EID, PIA
; APPLICANT: FARIS, MARY
 APPLICANT: HUBERT, RENE
  APPLICANT: GE, WANGMAO
  APPLICANT: JAKOBOVITZ, AYA
  TITLE OF INVENTION: NUCLEIC ACID AND CORRESPONDING PROTEIN ENTITLED
  TITLE OF INVENTION: 193P1E1B USEFUL IN TREATMENT AND DETECTION OF CANCER
; FILE REFERENCE: 51158-20063.00
; CURRENT APPLICATION NUMBER: US/10/013,312
; CURRENT FILING DATE: 2002-05-30
; NUMBER OF SEQ ID NOS: 3005
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; SOFTWARE: PatentIn Ver. 2.1
; SEQ ID NO 196
   LENGTH: 10
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   ORGANISM: Homo sapiens
US-10-013-312-196
                         31.5%; Score 28; DB 12; Length 10;
 Query Match
 Best Local Similarity 57.1%; Pred. No. 8.7e+02;
           4; Conservative 2; Mismatches 1; Indels
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                                                                           0;
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Qу
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Db
           2 TPSPIIQ 8
RESULT 26
US-10-013-312-459
; Sequence 459, Application US/10013312
; Publication No. US20030223990A1
; GENERAL INFORMATION:
; APPLICANT: RAITANO, ARTHUR
; APPLICANT: CHALLITA-EID, PIA
  APPLICANT: FARIS, MARY
  APPLICANT: HUBERT, RENE
  APPLICANT: GE, WANGMAO
  APPLICANT: JAKOBOVITZ, AYA
  TITLE OF INVENTION: NUCLEIC ACID AND CORRESPONDING PROTEIN ENTITLED
 TITLE OF INVENTION: 193P1E1B USEFUL IN TREATMENT AND DETECTION OF CANCER
; FILE REFERENCE: 51158-20063.00
; CURRENT APPLICATION NUMBER: US/10/013,312
; CURRENT FILING DATE: 2002-05-30
; NUMBER OF SEQ ID NOS: 3005
  SOFTWARE: PatentIn Ver. 2.1
; SEO ID NO 459
   LENGTH: 10
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   ORGANISM: Homo sapiens
US-10-013-312-459
                         31.5%; Score 28; DB 12; Length 10;
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Qу
             11 1::1
           2 TPSPIIQ 8
RESULT 27
US-10-013-312-560
; Sequence 560, Application US/10013312
; Publication No. US20030223990A1
; GENERAL INFORMATION:
; APPLICANT: RAITANO, ARTHUR
; APPLICANT: CHALLITA-EID, PIA
; APPLICANT: FARIS, MARY
```

```
; APPLICANT: HUBERT, RENE
; APPLICANT: GE, WANGMAO
; APPLICANT: JAKOBOVITZ, AYA
  TITLE OF INVENTION: NUCLEIC ACID AND CORRESPONDING PROTEIN ENTITLED
  TITLE OF INVENTION: 193P1E1B USEFUL IN TREATMENT AND DETECTION OF CANCER
  FILE REFERENCE: 51158-20063.00
  CURRENT APPLICATION NUMBER: US/10/013,312
; CURRENT FILING DATE: 2002-05-30
; NUMBER OF SEO ID NOS: 3005
; SOFTWARE: PatentIn Ver. 2.1
; SEQ ID NO 560
  LENGTH: 10
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  ORGANISM: Homo sapiens
US-10-013-312-560
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           3 TPQPLLQ 9
Qу
            2 TPSPIIQ 8
Db
RESULT 28
US-10-013-312-690
; Sequence 690, Application US/10013312
; Publication No. US20030223990A1
; GENERAL INFORMATION:
; APPLICANT: RAITANO, ARTHUR
; APPLICANT: CHALLITA-EID, PIA
; APPLICANT: FARIS, MARY
  APPLICANT: HUBERT, RENE
  APPLICANT: GE, WANGMAO
;
  APPLICANT: JAKOBOVITZ, AYA
  TITLE OF INVENTION: NUCLEIC ACID AND CORRESPONDING PROTEIN ENTITLED
  TITLE OF INVENTION: 193P1E1B USEFUL IN TREATMENT AND DETECTION OF CANCER
; FILE REFERENCE: 51158-20063.00
; CURRENT APPLICATION NUMBER: US/10/013,312
  CURRENT FILING DATE: 2002-05-30
; NUMBER OF SEQ ID NOS: 3005
  SOFTWARE: PatentIn Ver. 2.1
; SEQ ID NO 690
   LENGTH: 10
   TYPE: PRT
   ORGANISM: Homo sapiens
US-10-013-312-690
 Query Match
                        31.5%; Score 28; DB 12; Length 10;
 Best Local Similarity 57.1%; Pred. No. 8.7e+02;
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Qу
           3 TPQPLLQ 9
             11 1::1
Db
           2 TPSPIIQ 8
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RESULT 29
US-10-013-312-2072
; Sequence 2072, Application US/10013312
; Publication No. US20030223990A1
; GENERAL INFORMATION:
; APPLICANT: RAITANO, ARTHUR
; APPLICANT: CHALLITA-EID, PIA
; APPLICANT: FARIS, MARY
 APPLICANT: HUBERT, RENE
 APPLICANT: GE, WANGMAO
  APPLICANT: JAKOBOVITZ, AYA
  TITLE OF INVENTION: NUCLEIC ACID AND CORRESPONDING PROTEIN ENTITLED
; TITLE OF INVENTION: 193P1E1B USEFUL IN TREATMENT AND DETECTION OF CANCER
; FILE REFERENCE: 51158-20063.00
; CURRENT APPLICATION NUMBER: US/10/013,312
; CURRENT FILING DATE: 2002-05-30
; NUMBER OF SEQ ID NOS: 3005
  SOFTWARE: PatentIn Ver. 2.1
; SEQ ID NO 2072
   LENGTH: 10
   TYPE: PRT
   ORGANISM: Homo sapiens
US-10-013-312-2072
  Query Match
                         31.5%; Score 28; DB 12; Length 10;
  Best Local Similarity 57.1%; Pred. No. 8.7e+02;
          4; Conservative 2; Mismatches 1; Indels
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           3 TPQPLLQ 9
Qу
             11 1::1
           2 TPSPIIQ 8
RESULT 30
US-10-013-312-2241
; Sequence 2241, Application US/10013312
; Publication No. US20030223990A1
; GENERAL INFORMATION:
; APPLICANT: RAITANO, ARTHUR
  APPLICANT: CHALLITA-EID, PIA
  APPLICANT: FARIS, MARY
  APPLICANT: HUBERT, RENE
  APPLICANT: GE, WANGMAO
  APPLICANT: JAKOBOVITZ, AYA
  TITLE OF INVENTION: NUCLEIC ACID AND CORRESPONDING PROTEIN ENTITLED
  TITLE OF INVENTION: 193P1E1B USEFUL IN TREATMENT AND DETECTION OF CANCER
 FILE REFERENCE: 51158-20063.00
  CURRENT APPLICATION NUMBER: US/10/013,312
  CURRENT FILING DATE: 2002-05-30
  NUMBER OF SEQ ID NOS: 3005
 SOFTWARE: PatentIn Ver. 2.1
; SEQ ID NO 2241
   LENGTH: 10
    TYPE: PRT
    ORGANISM: Homo sapiens
US-10-013-312-2241
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31.5%; Score 28; DB 12; Length 10;
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           3 TPQPLLQ 9
Qy
             11 1::1
           2 TPSPIIQ 8
Db
RESULT 31
US-10-013-312-2432
; Sequence 2432, Application US/10013312
; Publication No. US20030223990A1
; GENERAL INFORMATION:
; APPLICANT: RAITANO, ARTHUR
; APPLICANT: CHALLITA-EID, PIA
  APPLICANT: FARIS, MARY
  APPLICANT: HUBERT, RENE
  APPLICANT: GE, WANGMAO
  APPLICANT: JAKOBOVITZ, AYA
  TITLE OF INVENTION: NUCLEIC ACID AND CORRESPONDING PROTEIN ENTITLED
  TITLE OF INVENTION: 193P1E1B USEFUL IN TREATMENT AND DETECTION OF CANCER
 FILE REFERENCE: 51158-20063.00
  CURRENT APPLICATION NUMBER: US/10/013,312
  CURRENT FILING DATE: 2002-05-30
  NUMBER OF SEQ ID NOS: 3005
  SOFTWARE: PatentIn Ver. 2.1
; SEQ ID NO 2432
   LENGTH: 10
    TYPE: PRT
    ORGANISM: Homo sapiens
US-10-013-312-2432
                         31.5%; Score 28; DB 12; Length 10;
  Query Match
  Best Local Similarity 57.1%; Pred. No. 8.7e+02;
          4; Conservative
                                2; Mismatches 1; Indels 0; Gaps
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  Matches
           3 TPQPLLQ 9
Qу
             11 1::1
Db
           1 TPSPIIQ 7
RESULT 32
US-10-013-312-2450
; Sequence 2450, Application US/10013312
; Publication No. US20030223990A1
; GENERAL INFORMATION:
  APPLICANT: RAITANO, ARTHUR
   APPLICANT: CHALLITA-EID, PIA
  APPLICANT: FARIS, MARY
  APPLICANT: HUBERT, RENE
  APPLICANT: GE, WANGMAO
  APPLICANT: JAKOBOVITZ, AYA
  TITLE OF INVENTION: NUCLEIC ACID AND CORRESPONDING PROTEIN ENTITLED
; TITLE OF INVENTION: 193P1E1B USEFUL IN TREATMENT AND DETECTION OF CANCER
; FILE REFERENCE: 51158-20063.00
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; CURRENT APPLICATION NUMBER: US/10/013,312
; CURRENT FILING DATE: 2002-05-30
; NUMBER OF SEQ ID NOS: 3005
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; SEQ ID NO 2450
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   ORGANISM: Homo sapiens
US-10-013-312-2450
                         31.5%; Score 28; DB 12; Length 10;
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Qу
             11 1::1
           2 TPSPIIQ 8
RESULT 33
US-10-234-816-33
; Sequence 33, Application US/10234816
; Publication No. US20030157514A1
; GENERAL INFORMATION:
; APPLICANT: Bristol-Myers Squibb Company
  TITLE OF INVENTION: POLYNUCLEOTIDE ENCODING A NOVEL PLECKSTRIN HOMOLOGY
DOMAIN AND PROLINE
; TITLE OF INVENTION: RICH DOMAIN CONTAINING ADAPTER PROTEIN, PMN29
; FILE REFERENCE: D0117 NP
; CURRENT APPLICATION NUMBER: US/10/234,816
; CURRENT FILING DATE: 2002-09-04
; NUMBER OF SEQ ID NOS: 118
 SOFTWARE: PatentIn version 3.1
; SEQ ID NO 33
   LENGTH: 13
   TYPE: PRT
   ORGANISM: Homo sapiens
US-10-234-816-33
                         31.5%; Score 28; DB 14; Length 13;
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 Best Local Similarity
                        57.1%; Pred. No. 1.2e+03;
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           4; Conservative
           2 QTPQPLL 8
Qу
             :||:||:
Db
           5 ETPRPLM 11
RESULT 34
US-10-185-425-24
; Sequence 24, Application US/10185425
; Publication No. US20030053984A1
; GENERAL INFORMATION:
; APPLICANT: Apotech Research and Development Ltd.
; APPLICANT: Tschopp, Jurg
; APPLICANT: Schneider, Pascal
; APPLICANT: Holler, Nils
```

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TITLE OF INVENTION: A Bimer or an Oligomer of a Dimer, Trimer, Quadromer, or
Pentamer of
  TITLE OF INVENTION: Recombinant Fusion Proteins
  FILE REFERENCE: 1708.001US1
  CURRENT APPLICATION NUMBER: US/10/185,425
  CURRENT FILING DATE: 2002-06-28
  PRIOR APPLICATION NUMBER: PCT/EP00/13032
  PRIOR FILING DATE: 2000-12-20
  PRIOR APPLICATION NUMBER: DE 199 63 859.4
  PRIOR FILING DATE: 1999-12-30
  NUMBER OF SEQ ID NOS: 24
  SOFTWARE: FastSEQ for Windows Version 4.0
; SEQ ID NO 24
   LENGTH: 14
   TYPE: PRT
   ORGANISM: Artificial Sequence
   FEATURE:
   OTHER INFORMATION: Linker
US-10-185-425-24
                         31.5%; Score 28; DB 14; Length 14;
  Query Match
 Best Local Similarity 50.0%; Pred. No. 1.2e+03;
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Qу
           4 POPLLOVMMEPQ 15
             111 1 11:
           3 РОРКРОРКРЕРЕ 14
RESULT 35
US-10-013-312-2620
; Sequence 2620, Application US/10013312
; Publication No. US20030223990A1
; GENERAL INFORMATION:
; APPLICANT: RAITANO, ARTHUR
; APPLICANT: CHALLITA-EID, PIA
; APPLICANT: FARIS, MARY
  APPLICANT: HUBERT, RENE
  APPLICANT: GE, WANGMAO
  APPLICANT: JAKOBOVITZ, AYA
  TITLE OF INVENTION: NUCLEIC ACID AND CORRESPONDING PROTEIN ENTITLED
  TITLE OF INVENTION: 193P1E1B USEFUL IN TREATMENT AND DETECTION OF CANCER
  FILE REFERENCE: 51158-20063.00
  CURRENT APPLICATION NUMBER: US/10/013,312
  CURRENT FILING DATE: 2002-05-30
; NUMBER OF SEQ ID NOS: 3005
  SOFTWARE: PatentIn Ver. 2.1
; SEQ ID NO 2620
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    ORGANISM: Homo sapiens
US-10-013-312-2620
  Query Match
                         31.5%; Score 28; DB 12; Length 15;
  Best Local Similarity 57.1%; Pred. No. 1.3e+03;
  Matches 4; Conservative 2; Mismatches 1; Indels
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            7 TPSPIIQ 13
RESULT 36
US-10-013-312-2653
; Sequence 2653, Application US/10013312
; Publication No. US20030223990A1
; GENERAL INFORMATION:
; APPLICANT: RAITANO, ARTHUR
; APPLICANT: CHALLITA-EID, PIA
; APPLICANT: FARIS, MARY
; APPLICANT: HUBERT, RENE
; APPLICANT: GE, WANGMAO
 APPLICANT: JAKOBOVITZ, AYA
  TITLE OF INVENTION: NUCLEIC ACID AND CORRESPONDING PROTEIN ENTITLED
  TITLE OF INVENTION: 193P1E1B USEFUL IN TREATMENT AND DETECTION OF CANCER
  FILE REFERENCE: 51158-20063.00
  CURRENT APPLICATION NUMBER: US/10/013,312
  CURRENT FILING DATE: 2002-05-30
; NUMBER OF SEQ ID NOS: 3005
 SOFTWARE: PatentIn Ver. 2.1
; SEQ ID NO 2653
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   ORGANISM: Homo sapiens
US-10-013-312-2653
                         31.5%; Score 28; DB 12; Length 15;
  Query Match
  Best Local Similarity 57.1%; Pred. No. 1.3e+03;
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Qy
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            1 TPSPIIQ 7
Db
RESULT 37
US-10-013-312-2691
; Sequence 2691, Application US/10013312
; Publication No. US20030223990A1
; GENERAL INFORMATION:
; APPLICANT: RAITANO, ARTHUR
; APPLICANT: CHALLITA-EID, PIA
; APPLICANT: FARIS, MARY
; APPLICANT: HUBERT, RENE
  APPLICANT: GE, WANGMAO
  APPLICANT: JAKOBOVITZ, AYA
   TITLE OF INVENTION: NUCLEIC ACID AND CORRESPONDING PROTEIN ENTITLED
   TITLE OF INVENTION: 193P1E1B USEFUL IN TREATMENT AND DETECTION OF CANCER
  FILE REFERENCE: 51158-20063.00
; CURRENT APPLICATION NUMBER: US/10/013,312
; CURRENT FILING DATE: 2002-05-30
; NUMBER OF SEQ ID NOS: 3005
; SOFTWARE: PatentIn Ver. 2.1
; SEQ ID NO 2691
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LENGTH: 15
   TYPE: PRT
  ORGANISM: Homo sapiens
US-10-013-312-2691
                        31.5%; Score 28; DB 12; Length 15;
 Query Match
 Best Local Similarity 57.1%; Pred. No. 1.3e+03;
 Matches 4; Conservative 2; Mismatches 1; Indels 0; Gaps
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           3 TPQPLLQ 9
Qу
             | | | | :: |
Db
           8 TPSPIIQ 14
RESULT 38
US-10-013-312-2692
; Sequence 2692, Application US/10013312
; Publication No. US20030223990A1
; GENERAL INFORMATION:
; APPLICANT: RAITANO, ARTHUR
; APPLICANT: CHALLITA-EID, PIA
; APPLICANT: FARIS, MARY
; APPLICANT: HUBERT, RENE
; APPLICANT: GE, WANGMAO
; APPLICANT: JAKOBOVITZ, AYA
  TITLE OF INVENTION: NUCLEIC ACID AND CORRESPONDING PROTEIN ENTITLED
; TITLE OF INVENTION: 193P1E1B USEFUL IN TREATMENT AND DETECTION OF CANCER
; FILE REFERENCE: 51158-20063.00
; CURRENT APPLICATION NUMBER: US/10/013,312
; CURRENT FILING DATE: 2002-05-30
; NUMBER OF SEO ID NOS: 3005
; SOFTWARE: PatentIn Ver. 2.1
; SEQ ID NO 2692
   LENGTH: 15
   TYPE: PRT
   ORGANISM: Homo sapiens
US-10-013-312-2692
                        31.5%; Score 28; DB 12; Length 15;
  Query Match
  Best Local Similarity 57.1%; Pred. No. 1.3e+03;
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Qу
             | | | ::|
Db
           6 TPSPIIQ 12
RESULT 39
US-10-013-312-2781
; Sequence 2781, Application US/10013312
; Publication No. US20030223990A1
; GENERAL INFORMATION:
; APPLICANT: RAITANO, ARTHUR
; APPLICANT: CHALLITA-EID, PIA
; APPLICANT: FARIS, MARY
; APPLICANT: HUBERT, RENE
; APPLICANT: GE, WANGMAO
```

```
; APPLICANT: JAKOBOVITZ, AYA
  TITLE OF INVENTION: NUCLEIC ACID AND CORRESPONDING PROTEIN ENTITLED.
; TITLE OF INVENTION: 193P1E1B USEFUL IN TREATMENT AND DETECTION OF CANCER
; FILE REFERENCE: 51158-20063.00
  CURRENT APPLICATION NUMBER: US/10/013,312
; CURRENT FILING DATE: 2002-05-30
; NUMBER OF SEQ ID NOS: 3005
; SOFTWARE: PatentIn Ver. 2.1
; SEO ID NO 2781
  LENGTH: 15
   TYPE: PRT
   ORGANISM: Homo sapiens
US-10-013-312-2781
  Query Match
                       31.5%; Score 28; DB 12; Length 15;
 Best Local Similarity 57.1%; Pred. No. 1.3e+03;
          4; Conservative 2; Mismatches 1; Indels 0; Gaps
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                                                                         0;
           3 TPQPLLQ 9
Qу
            7 TPSPIIQ 13
Db
RESULT 40
US-10-013-312-2808
; Sequence 2808, Application US/10013312
; Publication No. US20030223990A1
; GENERAL INFORMATION:
; APPLICANT: RAITANO, ARTHUR
; APPLICANT: CHALLITA-EID, PIA
; APPLICANT: FARIS, MARY
; APPLICANT: HUBERT, RENE
; APPLICANT: GE, WANGMAO
 APPLICANT: JAKOBOVITZ, AYA
  TITLE OF INVENTION: NUCLEIC ACID AND CORRESPONDING PROTEIN ENTITLED
 TITLE OF INVENTION: 193P1E1B USEFUL IN TREATMENT AND DETECTION OF CANCER
; FILE REFERENCE: 51158-20063.00
; CURRENT APPLICATION NUMBER: US/10/013,312
; CURRENT FILING DATE: 2002-05-30
; NUMBER OF SEQ ID NOS: 3005
  SOFTWARE: PatentIn Ver. 2.1
; SEQ ID NO 2808
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   TYPE: PRT
   ORGANISM: Homo sapiens
US-10-013-312-2808
                       31.5%; Score 28; DB 12; Length 15;
  Query Match
  Best Local Similarity 57.1%; Pred. No. 1.3e+03;
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Qу
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            11 1::1
Db
           2 TPSPIIO 8
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RESULT 41

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US-10-013-312-2824
; Sequence 2824, Application US/10013312
; Publication No. US20030223990A1
; GENERAL INFORMATION:
; APPLICANT: RAITANO, ARTHUR
  APPLICANT: CHALLITA-EID, PIA
  APPLICANT: FARIS, MARY
; APPLICANT: HUBERT, RENE
; APPLICANT: GE, WANGMAO
  APPLICANT: JAKOBOVITZ, AYA
  TITLE OF INVENTION: NUCLEIC ACID AND CORRESPONDING PROTEIN ENTITLED
  TITLE OF INVENTION: 193P1E1B USEFUL IN TREATMENT AND DETECTION OF CANCER
  FILE REFERENCE: 51158-20063.00
; CURRENT APPLICATION NUMBER: US/10/013,312
; CURRENT FILING DATE: 2002-05-30
; NUMBER OF SEQ ID NOS: 3005
  SOFTWARE: PatentIn Ver. 2.1
; SEQ ID NO 2824
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   TYPE: PRT
   ORGANISM: Homo sapiens
US-10-013-312-2824
 Query Match
                         31.5%; Score 28; DB 12; Length 15;
 Best Local Similarity 57.1%; Pred. No. 1.3e+03;
           4; Conservative 2; Mismatches 1; Indels
                                                                0; Gaps
 Matches
                                                                            0;
           3 TPQPLLQ 9
Qy
             11 1::1
Db
           6 TPSPIIQ 12
RESULT 42
US-10-161-791-195
; Sequence 195, Application US/10161791
; Publication No. US20030186863A1
  GENERAL INFORMATION:
    APPLICANT: SPARKS, Andrew B.
    APPLICANT: KAY, Brian K.
    APPLICANT: THORN, Judith M.
    APPLICANT: QUILLIAM, Lawrence A.
    APPLICANT: DER, Channing J.
    APPLICANT: FOWLKES, Dana M.
    APPLICANT: RIDER, James E.
    TITLE OF INVENTION: SH3 BINDING PEPTIDES AND METHODS OF
    TITLE OF INVENTION: ISOLATING AND USING SAME
    NUMBER OF SEQUENCES: 467
    CORRESPONDENCE ADDRESS:
      ADDRESSEE: Pennie & Edmonds
      STREET: 1155 Avenue of the Americas
      CITY: New York
      STATE: New York
      COUNTRY: U.S.A.
      ZIP: 10036-2711
    COMPUTER READABLE FORM:
      MEDIUM TYPE: Floppy disk
      COMPUTER: IBM PC compatible
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OPERATING SYSTEM: PC-DOS/MS-DOS
      SOFTWARE: PatentIn Release #1.0, Version #1.30
    CURRENT APPLICATION DATA:
      APPLICATION NUMBER: US/10/161,791
      FILING DATE:
      CLASSIFICATION:
;
    PRIOR APPLICATION DATA:
;
      APPLICATION NUMBER: US/08/602,999
      FILING DATE: 16-FEB-1996
    ATTORNEY/AGENT INFORMATION:
;
      NAME: Misrock, S. Leslie
      REGISTRATION NUMBER: 18,872
;
      REFERENCE/DOCKET NUMBER: 1101-202
    TELECOMMUNICATION INFORMATION:
      TELEPHONE: (212) 790-9090
      TELEFAX: (212) 869-9741/8864
      TELEX: 66141 PENNIE
  INFORMATION FOR SEQ ID NO: 195:
    SEQUENCE CHARACTERISTICS:
      LENGTH: 16 amino acids
      TYPE: amino acid
;
      TOPOLOGY: unknown
    MOLECULE TYPE: peptide
US-10-161-791-195
                         31.5%; Score 28; DB 14; Length 16;
 Query Match
  Best Local Similarity 41.7%; Pred. No. 1.4e+03;
           5; Conservative 2; Mismatches 5; Indels
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           3 TPQPLLQVMMEP 14
Qу
             11:1: 11
           4 TPRPAVPQRMNP 15
RESULT 43
US-10-319-315-49
; Sequence 49, Application US/10319315
; Publication No. US20030219774A1
; GENERAL INFORMATION:
  APPLICANT: Bristol-Myers Squibb Company
  TITLE OF INVENTION: NOVEL HUMAN NEUROTRANSMITTER TRANSPORTER
  FILE REFERENCE: D0205 NP
  CURRENT APPLICATION NUMBER: US/10/319,315
; CURRENT FILING DATE: 2002-12-13
; NUMBER OF SEQ ID NOS: 135
  SOFTWARE: PatentIn version 3.1
; SEQ ID NO 49
   LENGTH: 16
   TYPE: PRT
   ORGANISM: Homo sapiens
US-10-319-315-49
  Query Match
                         31.5%; Score 28; DB 15; Length 16;
  Best Local Similarity 50.0%; Pred. No. 1.4e+03;
                               2; Mismatches 3; Indels
 Matches
           5; Conservative
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Qу
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3 PKLDKMLDPQ 12
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.Db

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RESULT 44
US-09-991-225-68
; Sequence 68, Application US/09991225
; Publication No. US20030153063A1
; GENERAL INFORMATION:
; APPLICANT: Bristol-Myers Squibb Company
   TITLE OF INVENTION: A NOVEL HUMAN G-PROTEIN COUPLED RECEPTOR, HGPRBMY11,
EXPRESSED HIGHLY IN
; TITLE OF INVENTION: HEART AND VARIANTS THEREOF
 FILE REFERENCE: D0075.NP
; CURRENT APPLICATION NUMBER: US/09/991,225
; CURRENT FILING DATE: 2001-11-16
; PRIOR APPLICATION NUMBER: 60/249,613
 PRIOR FILING DATE: 2000-11-17
  PRIOR APPLICATION NUMBER: 60/257,611
  PRIOR FILING DATE: 2000-12-21
  PRIOR APPLICATION NUMBER: 60/305,818
  PRIOR FILING DATE: 2001-07-16
 NUMBER OF SEQ ID NOS: 81
  SOFTWARE: PatentIn version 3.0
; SEQ ID NO 68
   LENGTH: 14
    TYPE: PRT
    ORGANISM: Homo sapiens
US-09-991-225-68
  Query Match
                         30.9%; Score 27.5; DB 10; Length 14;
  Best Local Similarity 53.8%; Pred. No. 1.5e+03;
  Matches
           7; Conservative 1; Mismatches
                                                 4; Indels
                                                                 1; Gaps
                                                                             1;
Qv
            5 QPLLQVM-MEPQG 16
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            2 QPSISVSEMEPNG 14
Db
RESULT 45
US-10-369-405-68
; Sequence 68, Application US/10369405
; Publication No. US20030224400A1
; GENERAL INFORMATION:
; APPLICANT: Bristol-Myers Squibb Company
; TITLE OF INVENTION: NOVEL HUMAN G-PROTEIN COUPLED RECEPTOR, HGPRBMY11, AND
VARIANTS
; TITLE OF INVENTION: THEREOF
  FILE REFERENCE: D0075A CIP
  CURRENT APPLICATION NUMBER: US/10/369,405
  CURRENT FILING DATE: 2003-02-14
 PRIOR APPLICATION NUMBER: U.S. 60/249,613
 PRIOR FILING DATE: 2000-11-17
 PRIOR APPLICATION NUMBER: U.S. 09/991,225
; PRIOR FILING DATE: 2001-11-16
; PRIOR APPLICATION NUMBER: U.S. 60/257,611
; PRIOR FILING DATE: 2000-12-21
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; PRIOR APPLICATION NUMBER: U.S. 60/305,818
; PRIOR FILING DATE: 2001-07-16
; NUMBER OF SEQ ID NOS: 94
 SOFTWARE: PatentIn version 3.2
; SEQ ID NO 68
   LENGTH: 14
   TYPE: PRT
   ORGANISM: Homo sapiens
US-10-369-405-68
                         30.9%; Score 27.5; DB 12; Length 14;
  Query Match
  Best Local Similarity 53.8%; Pred. No. 1.5e+03;
           7; Conservative 1; Mismatches 4; Indels 1; Gaps
 Matches
                                                                          1;
Qy
           5 QPLLQVM-MEPQG 16
             Db
           2 QPSISVSEMEPNG 14
RESULT 46
US-09-984-056-95
; Sequence 95, Application US/09984056
; Patent No. US20020120106A1
; GENERAL INFORMATION:
; APPLICANT: BOGOCH, SAMUEL
  APPLICANT: BOGOCH, ELENORE S.
  TITLE OF INVENTION: ANTHRAX AND SMALL POX REPLIKINS AND METHODS OF USE
  FILE REFERENCE: 09425-46903
; CURRENT APPLICATION NUMBER: US/09/984,056
; CURRENT FILING DATE: 2001-10-26
  PRIOR APPLICATION NUMBER: 60/303,396
; PRIOR FILING DATE: 2001-07-09
; PRIOR APPLICATION NUMBER: 60/278,761
  PRIOR FILING DATE: 2001-03-27
  PRIOR APPLICATION NUMBER: 09/146,755
  PRIOR FILING DATE: 1998-09-04
  PRIOR APPLICATION NUMBER: 09/817,144
  PRIOR FILING DATE: 2001-03-27
  PRIOR APPLICATION NUMBER: 08/198,139
 PRIOR FILING DATE: 1994-02-17
; NUMBER OF SEQ ID NOS: 103
  SOFTWARE: PatentIn 2.1
; SEQ ID NO 95
   LENGTH: 18
   TYPE: PRT
   ORGANISM: Bacillus anthracis
US-09-984-056-95
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  Query Match
  Best Local Similarity 41.2%; Pred. No. 1.9e+03;
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                        11: 1
           3 LQIPPPI----EPKKD 14
Db
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RESULT 47
US-10-105-232-95
; Sequence 95, Application US/10105232
; Publication No. US20030180328A1
; GENERAL INFORMATION:
  APPLICANT: BOGOCH, SAMUEL
 APPLICANT: BOGOCH, ELENORE S.
; TITLE OF INVENTION: REPLIKIN PEPTIDES IN RAPID REPLICATION OF GLIOMA CELLS
  TITLE OF INVENTION: AND IN INFLUENZA EPIDEMICS
; FILE REFERENCE: 09425-46904
  CURRENT APPLICATION NUMBER: US/10/105,232
  CURRENT FILING DATE: 2002-03-26
  PRIOR APPLICATION NUMBER: 60/303,396
; PRIOR FILING DATE: 2001-07-09
; PRIOR APPLICATION NUMBER: 60/278,761
; PRIOR FILING DATE: 2001-03-27
; PRIOR APPLICATION NUMBER: 09/146,755
; PRIOR FILING DATE: 1998-09-04
  PRIOR APPLICATION NUMBER: 09/817,144
  PRIOR FILING DATE: 2001-03-27
; PRIOR APPLICATION NUMBER: 08/198,139
; PRIOR FILING DATE: 1994-02-17
; NUMBER OF SEQ ID NOS: 535
  SOFTWARE: PatentIn 2.1
; SEQ ID NO 95
   LENGTH: 18
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   ORGANISM: Bacillus anthracis
US-10-105-232-95
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  Best Local Similarity 41.2%; Pred. No. 1.9e+03;
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Qy
           1 LQTPQPLLQVMMEPQGD 17
                        ||:|
             | | | | :
            3 LQIPPPI----EPKKD 14
RESULT 48
US-10-185-815-89
; Sequence 89, Application US/10185815
; Publication No. US20030096354A1
; GENERAL INFORMATION:
; APPLICANT: Elan Corporation, plc
; APPLICANT: O'Mahony, Daniel
; APPLICANT: Lambkin, Imelda
; APPLICANT: Higgins, Lisa
  TITLE OF INVENTION: Peyer's Patch And/Or M-Cell Targeting Ligands
  FILE REFERENCE: E1067-20093
  CURRENT APPLICATION NUMBER: US/10/185,815
  CURRENT FILING DATE: 2002-10-09
; PRIOR APPLICATION NUMBER: 60/302,591
; PRIOR FILING DATE: 2001-07-02
; NUMBER OF SEQ ID NOS: 99
; SOFTWARE: PatentIn version 3.1
; SEQ ID NO 89
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LENGTH: 10
   TYPE: PRT
   ORGANISM: Artificial Sequence
   FEATURE:
   OTHER INFORMATION: Segment of CAPHI beta casein precursor
US-10-185-815-89
                         30.3%; Score 27; DB 14; Length 10;
  Query Match
  Best Local Similarity 83.3%; Pred. No. 1.2e+03;
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 Matches
           2 QTPQPL 7
Qу
             1 1111
Dh
           5 QPPQPL 10
RESULT 49
US-09-954-385-136
; Sequence 136, Application US/09954385
; Publication No. US20030100467A1
; GENERAL INFORMATION:
; APPLICANT: Aehle, Wolfgang
; APPLICANT: Baldwin, Toby L.
; APPLICANT: Van Gastel, Franciscus J.C.
  APPLICANT: Janssen, Giselle G.
  APPLICANT: Murray, Christopher J.
; APPLICANT: Wang, Huaming
; APPLICANT: Winetzky, Deborah S.
  TITLE OF INVENTION: Binding Phenol Oxidizing Enzyme-peptide
; TITLE OF INVENTION: Complexes
; FILE REFERENCE: GC690
; CURRENT APPLICATION NUMBER: US/09/954,385
; CURRENT FILING DATE: 2001-09-12
  NUMBER OF SEQ ID NOS: 433
  SOFTWARE: FastSEQ for Windows Version 4.0
; SEQ ID NO 136
   LENGTH: 12
   TYPE: PRT
   ORGANISM: Artificial Sequence
   FEATURE:
    OTHER INFORMATION: binding peptide
US-09-954-385-136
                         30.3%; Score 27; DB 10; Length 12;
  Query Match
                         71.4%; Pred. No. 1.5e+03;
  Best Local Similarity
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           3 TPQPLLQ 9
Qу
              11 111:
Db
            6 TPPPLLR 12
RESULT 50
US-09-749-873-125
; Sequence 125, Application US/09749873
; Publication No. US20030023045A1
   GENERAL INFORMATION:
```

```
APPLICANT: OHTOMO, Toshihiko
                    SATO, Koh
                    TSUCHIYA, Masayuki
         TITLE OF INVENTION: RESHAPED HUMAN ANTIBODY TO HUMAN
                             MEDULLOBLASTOMA CELLS
         NUMBER OF SEQUENCES: 132
         CORRESPONDENCE ADDRESS:
              ADDRESSEE: Foley & Lardner
              STREET: 3000 K Street, N.W., Suite 500
              CITY: Washington
              STATE: D.C.
              COUNTRY: USA
              ZIP: 20007-5109
        COMPUTER READABLE FORM:
              MEDIUM TYPE: Floppy disk
              COMPUTER: IBM PC compatible
              OPERATING SYSTEM: PC-DOS/MS-DOS
              SOFTWARE: PatentIn Release #1.0, Version #1.30
         CURRENT APPLICATION DATA:
              APPLICATION NUMBER: US/09/749,873
              FILING DATE: 29-Dec-2000
              CLASSIFICATION: <Unknown>
         PRIOR APPLICATION DATA:
              APPLICATION NUMBER: 08/646,265
              FILING DATE: 1996-09-09
              APPLICATION NUMBER: JP 5-291078
              FILING DATE: 19-NOV-1993
        ATTORNEY/AGENT INFORMATION:
              NAME: WEGNER, Harold C.
              REGISTRATION NUMBER: 25,258
              REFERENCE/DOCKET NUMBER: 53466/184
         TELECOMMUNICATION INFORMATION:
              TELEPHONE: (202) 672-5300
              TELEFAX: (202) 672-5399
              TELEX: 904136
    INFORMATION FOR SEQ ID NO: 125:
        SEQUENCE CHARACTERISTICS:
              LENGTH: 15 amino acids
              TYPE: amino acid
              TOPOLOGY: linear
         MOLECULE TYPE: protein
         SEQUENCE DESCRIPTION: SEQ ID NO: 125:
US-09-749-873-125
                          30.3%; Score 27; DB 10; Length 15;
  Query Match
  Best Local Similarity 57.1%; Pred. No. 1.9e+03;
           4; Conservative 3; Mismatches 0; Indels 0; Gaps
  Matches
                                                                             0;
           2 QTPQPLL 8
Qу
              1:1:11:
            8 OSPKPLI 14
Search completed: July 4, 2004, 05:12:23
Job time : 35.3657 secs
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## GenCore version 5.1.6 Copyright (c) 1993 - 2004 Compugen Ltd.

OM protein - protein search, using sw model

July 4, 2004, 04:33:51; Search time 28.9254 Seconds Run on:

(without alignments)

185.436 Million cell updates/sec

Title:

US-09-641-802-2

Perfect score: 89

Sequence: 1 LQTPQPLLQVMMEPQGD 17

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

1017041 seqs, 315518202 residues Searched:

Total number of hits satisfying chosen parameters: 5664

Minimum DB seq length: 7 Maximum DB seq length: 18

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 100 summaries

SPTREMBL 25:\* Database :

> 1: sp archea:\* 2: sp bacteria:\* 3: sp fungi:\* 4: sp human:\*

5: sp\_invertebrate:\*

6: sp\_mammal:\* 7: sp mhc:\*

8: sp organelle:\* 9: sp\_phage:\* 10: sp\_plant:\* 11: sp\_rodent:\*

12: sp\_virus:\*
13: sp\_vertebrate:\*

14: sp\_unclassified:\*

15: sp\_rvirus:\* 16: sp bacteriap:\* 17: sp archeap:\*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

## SUMMARIES

ક Result Query

Description

No. Score Match Length DB ID

1	32	36.0	18	5	Q9TWM9	Q9twm9 tetrahymena
2	28	31.5	17	10		Q7y0i3 zea mays su
3	27	30.3	16	8	Q8MC25	Q8mc25 sonneratia
4	27	30.3	16	8	Q8MC39	Q8mc39 ammannia ba
5	27	30.3	16	8	Q8LVE1	Q81ve1 punica gran
6	27	30.3	16	8	Q8MC53	Q8mc53 woodfordia
7	27	30.3	16	8	Q8MC17	Q8mc17 ludwigia hy
8	27	30.3	16	8	Q8LVE2	Q81ve2 lythrum sal
9	27	30.3	16	8	Q8LVE0	Q81ve0 trapa maxim
10	27	30.3	16	8	Q8MET2	Q8met2 saxifraga s
11	27	30.3	16	8	Q8MC45	Q8mc45 decodon ver
12	27	30.3	16	8	Q8MC21	Q8mc21 combretum w
13	27	30.3	16	8	Q8MC51	Q8mc51 cuphea lanc
14	27	30.3	16	8	Q8MC83	Q8mc33 rotala indi
15	27	30.3	16	8	Q8MC19	Q8mc19 quisqualis
16	27	30.3	16	8	Q8MC49	Q8mc49 pemphis aci
17	27	30.3	16	8	Q8MC27	Q8mc27 lagerstroem
18	27	30.3	16	8	Q8MET0	Q8met0 cercidiphyl
19	27	30.3	16	8	Q8MC23	Q8mc23 sonneratia
20	27	30.3	16	8	Q8MC41	Q8mc41 nesaea lued
21	27	30.3	16	8	Q8MC15	Q8mc15 fuchsia hyb
22	27	30.3	16	8	Q8MC43	Q8mc43 sonneratia
23	27	30.3	16	8	Q8MC47	Q8mc47 peplis port
24	27	30.3	16	8	Q8MC37	Q8mc37 lawsonia in
25	27	30.3	16	8	Q8MC29	Q8mc29 duabanga gr
26	27	30.3	16	8	Q8MC31	Q8mc31 lagerstroem
27	27	30.3	16	8	Q8MES8	Q8mes8 daphniphyll
28	27	30.3	16	8	Q8MC35	Q8mc35 heimia myrt
29	27	30.3	17	8	Q9XQN9	Q9xqn9 sinapis alb
30	26	29.2	17	13	~	Q90xe2 gallus gall
31	25	28.1	16	4	Q9UCH1	Q9uch1 homo sapien
32	25	28.1	16	6	Q9TRB4	Q9trb4 bos taurus
33	25	28.1	17	2	Q9APT4	Q9apt4 pseudomonas
34	25	28.1	18	2	Q9R5F6	Q9r5f6 helicobacte
35	23	25.8	11	2	Q44237	Q44237 anabaena sp
36 37	23 23	25.8	13	4	Q9UEE2	Q9uee2 homo sapien
3 <i>1</i> 38	23 23	25.8 25.8	16 16	4 10	Q96RT5 Q84RM4	Q96rt5 homo sapien Q84rm4 boea crassi
39	23	25.8	17	2	Q7X3X0	Q7x3x0 streptomyce
40	23	25.8	18	7	Q7X3X0 Q30216	Q30216 homo sapien
41	22	24.7	12	11		Q8cg32 rattus sp.
42	22	24.7	13	11		088176 mus musculu
43	22	24.7	13	11		Q8ciw3 mus musculu
44	22	24.7	16	4	Q9NNZ2	Q9nnz2 homo sapien
45	22	24.7	18	16		Q81cs8 bacillus ce
46	21	23.6	8	8	Q9T2Y3	Q9t2y3 begonia for
47	21	23.6	10	11		Q7tsc5 mus musculu
48	21	23.6	15	2	Q9X637	Q9x637 klebsiella
49	21	23.6	15	2	Q9X635	Q9x635 escherichia
50	21	23.6	16	4	Q9UQS4	Q9uqs4 homo sapien
51	21	23.6	16	15		Q7sm54 human t-lym
52	21	23.6	17	12		Q8v9k6 human papil
53 <sup>-</sup>	21	23.6	17	15		Q9e124 human immun
54	21	23.6	18	2	Q9R5S8	Q9r5s8 treponema d
55	20	22.5	8	2	Q8GMM5	Q8gmm5 acinetobact
56	20	22.5	10	4	Q9UNF2	Q9unf2 homo sapien
57	20	22.5	13	12		Q81769 hepatitis c

58	20	22.5	15	7 Q9TNQ1	Q9tnq1 mus sp. bet
59	20	22.5	15	10 Q9S8N8	Q9s8n8 hordeum vul
60	20	22.5	15	11 Q9QV72	Q9qv72 mus sp. int
61	20	22.5	16	5 Q9TWK1	Q9twk1 mytilus edu
62	20	22.5	16	6 Q9TR99	Q9tr99 canis famil
63	20	22.5	16	8 Q9T2R0	Q9t2r0 solanum tub
64	20	22.5	16	16 Q9KDQ5	Q9kdq5 bacillus ha
65	20	22.5	17	11 Q9JLA7	Q9jla7 mus musculu
66	20	22.5	18	3 Q9UR73	Q9ur73 trichoderma
67	20	22.5	18	4 Q9UCF9	Q9ucf9 homo sapien
68	20	22.5	18	6 097773	097773 cercopithec
69	20	22.5	18	6 Q7YRH7	Q7yrh7 sus scrofa
70	20	22.5	18	11 Q7TNW8	Q7tnw8 rattus norv
71	20	22.5	18	11 Q7TNW7	Q7tnw7 mus musculu
72	19.5	21.9	15	10 P83184	P83184 triticum ae
73	19	21.3	10	12 Q9Q0W9	Q9q0w9 polyomaviru
74	19	21.3	10	12 Q8JV70	Q8jv70 polyomaviru
75	19	21.3	10	12 Q9Q0W1	Q9q0w1 polyomaviru
76	19	21.3	10	12 Q8JV68	Q8jv68 polyomaviru
77	19	21.3	10	12 Q9Q0V9	Q9q0v9 polyomaviru
78	19	21.3	10	12 Q9Q0W7	Q9q0w7 polyomaviru
79	19	21.3	10	12 Q8JV66	Q8jv66 polyomaviru
80	19	21.3	10	12 Q9Q0V7	Q9q0v7 polyomaviru
81	19	21.3	10	12 Q8JV82	Q8jv82 polyomaviru
82	19	21.3	10	12 Q8JV76	Q8jv76 polyomaviru
83	19	21.3	10	12 Q8JV74	Q8jv74 polyomaviru
84	19	21.3	10	12 Q9Q0X7	Q9q0x7 polyomaviru
85	19	21.3	10	12 Q9Q0W5	Q9q0w5 polyomaviru
86	19	21.3	10	12 Q9Q0X3	Q9q0x3 polyomaviru
87	19	21.3	10	12 Q9Q0X5	Q9q0x5 polyomaviru
88	19	21.3	10	12 Q9Q0W3	Q9q0w3 polyomaviru
89	19	21.3	10	12 Q8JV80	Q8jv80 polyomaviru
90	19	21.3	10	12 Q9Q0X1	Q9q0x1 polyomaviru
91	19	21.3	10	12 Q9Q0X9	Q9q0x9 polyomaviru
92	19	21.3	10	12 Q8JV72	Q8jv72 polyomaviru
93	19	21.3	11	4 Q9C057	Q9c057 homo sapien
94	19	21.3	11	12 Q80B26	Q80b26 hepatitis b
95	19	21.3	12	2 Q9R5F7	Q9r5f7 helicobacte
96	19	21.3	13	9 Q8HA05	Q8ha05 bacteriopha
97	19	21.3	14	2 Q8RK22	Q8rk22 pseudomonas
98	19	21.3	14	2 Q47599	Q47599 escherichia
99	19	21.3	14	15 Q8JDM3	Q8jdm3 human immun
100	19	21.3	14	15 Q8JDM7	Q8jdm7 human immun

## ALIGNMENTS

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                                  PRT;
                                          18 AA.
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    01-MAY-2000 (TrEMBLrel. 13, Created)
DT
     01-MAY-2000 (TrEMBLrel. 13, Last sequence update)
DT
DT
     01-JUN-2000 (TrEMBLrel. 14, Last annotation update)
DΕ
     NKTAG antigen (Fragment).
os
     Tetrahymena pyriformis.
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OC
     Eukaryota; Alveolata; Ciliophora; Oligohymenophorea; Hymenostomatida;
OC
     Tetrahymenina; Tetrahymena.
OX
     NCBI TaxID=5908;
RN
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RP
     SEQUENCE.
RX
    MEDLINE=94323706; PubMed=7519359;
     Leary J.H.3rd, Evans D.L., Jaso-Friedmann L.;
RA
RT
     "Partial amino acid sequence of a novel protozoan parasite antigen
     that inhibits non-specific cytotoxic cell activity.";
RT
     Scand. J. Immunol. 40:158-164(1994).
RL
              18 AA; 2082 MW; 13BCB737B1FF92A3 CRC64;
SQ
     SEQUENCE
  Query Match
                          36.0%; Score 32; DB 5; Length 18;
  Best Local Similarity
                          60.0%; Pred. No. 2.4e+02;
             6; Conservative
                              1; Mismatches
                                                 3; Indels
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                                                                             0;
Qу
           8 LQVMMEPQGD 17
             11: 1111
Db
           3 LQLPFNPQGD 12
RESULT 2
Q7Y0I3
ID
    Q7Y0I3
                PRELIMINARY;
                                   PRT;
                                          17 AA.
AC
     Q7Y0I3;
DT
     01-OCT-2003 (TrEMBLrel. 25, Created)
DT
     01-OCT-2003 (TrEMBLrel. 25, Last sequence update)
     01-OCT-2003 (TrEMBLrel. 25, Last annotation update)
DT
DE
     Isoamylase (Fragment).
GN
     SU1.
os
     Zea mays subsp. mays (maize).
OC
     Eukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta;
OC
     Spermatophyta; Magnoliophyta; Liliopsida; Poales; Poaceae;
OC
     PACCAD clade; Panicoideae; Andropogoneae; Zea.
OX
     NCBI TaxID=4578;
RN
    [1]
RP
     SEQUENCE FROM N.A.
RC
     STRAIN=cv. T232;
     Wilson L.M., Whitt S.R., Ibanez A.M., Rocheford T.R., Goodman M.M.,
RA
     Buckler E.S. IV.;
RA
RT
     "Dissection of maize starch production by candidate gene
RT
     association.";
     Submitted (MAY-2003) to the EMBL/GenBank/DDBJ databases.
RL
     EMBL; AY290391; AAP45416.1; -.
DR
FT
     NON TER
                  17
                        17
SQ
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               17 AA; 1810 MW; E2913BED794C0670 CRC64;
  Query Match
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  Best Local Similarity 50.0%; Pred. No. 1e+03;
            5; Conservative
                                3; Mismatches
                                                 2; Indels
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           1 LQTPQPLLQV 10
Qу
             :::::::
            8 VSSPRPLLAV 17
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Q8MC25
                                   PRT;
                                           16 AA.
ID
    Q8MC25
                 PRELIMINARY;
AC
    Q8MC25;
     01-OCT-2002 (TrEMBLrel. 22, Created)
DT
     01-OCT-2002 (TrEMBLrel. 22, Last sequence update)
DT
     01-JUN-2003 (TrEMBLrel. 24, Last annotation update)
DT
DE
     PsaA (Fragment).
GN
    PSAA.
OS
    Sonneratia apetala.
OG
    Chloroplast.
    Eukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta;
OC
     Spermatophyta; Magnoliophyta; eudicotyledons; core eudicots; rosids;
OC
    Myrtales; Lythraceae; Sonneratia.
OC
OX
    NCBI TaxID=122813;
RN
    [1]
    SEQUENCE FROM N.A.
RP
    STRAIN=S165;
RC
     Huang Y., Shi S.;
RA
     "Phylogenetics of Lythraceae sensu lato: A Preliminary Analysis Based
RT
     on Chloroplast rbcL Gene, psaA-ycf3 Spacer, and Nuclear rDNA Internal
RT
     Transcribed Spacer (ITS) Sequences.";
RT
     Int. J. Plant Sci. 163:215-225(2002).
RL
DR
    EMBL; AY035740; AAL14161.1; -.
    GO; GO:0009507; C:chloroplast; IEA.
DR
KW
    Chloroplast.
FT
     NON TER
                  16
                         16
                16 AA; 1895 MW; 517FE691B89355B9 CRC64;
SQ
     SEQUENCE
                          30.3%; Score 27; DB 8; Length 16;
  Query Match
                          15.4%; Pred. No. 1.4e+03;
  Best Local Similarity
                                                                              0;
           2: Conservative 10; Mismatches 1; Indels
                                                                  0; Gaps
Qу
            1 LQTPQPLLQVMME 13
              :::|:| ::::::
Db
            3 IRSPEPEVKILVD 15
RESULT 4
08MC39
     Q8MC39
                 PRELIMINARY;
                                   PRT:
                                           16 AA.
ID
AC
     Q8MC39;
     01-OCT-2002 (TrEMBLrel. 22, Created)
DT
     01-OCT-2002 (TrEMBLrel. 22, Last sequence update)
DT
     01-JUN-2003 (TrEMBLrel. 24, Last annotation update)
DT
DE
     PsaA (Fragment).
GN
     PSAA.
     Ammannia baccifera.
OS
     Chloroplast.
OG
     Eukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta;
OC
     Spermatophyta; Magnoliophyta; eudicotyledons; core eudicots; rosids;
OC
     Myrtales; Lythraceae; Ammannia.
OC
     NCBI TaxID=162022;
OX
RN
     [1]
RP
     SEOUENCE FROM N.A.
RC
     STRAIN=A610;
RA
     Huang Y., Shi S.;
RT
     "Phylogenetics of Lythraceae sensu lato: A Preliminary Analysis Based
```

```
on Chloroplast rbcL Gene, psaA-ycf3 Spacer, and Nuclear rDNA Internal
RT
    Transcribed Spacer (ITS) Sequences.";
RT
    Int. J. Plant Sci. 163:215-225(2002).
RL
    EMBL; AY035733; AAL14147.1; -.
DR
    GO; GO:0009507; C:chloroplast; IEA.
DR
KW
    Chloroplast.
    NON TER
                  16
FT
                         16
               16 AA; 1895 MW; 517FE691B89355B9 CRC64;
     SEQUENCE
SQ
                          30.3%; Score 27; DB 8; Length 16;
  Query Match
                         15.4%; Pred. No. 1.4e+03;
 Best Local Similarity
            2; Conservative 10; Mismatches
 Matches
                                                 1; Indels
                                                                 0; Gaps
Qу
           1 LQTPQPLLQVMME 13
             :::|:| ::::::
            3 IRSPEPEVKILVD 15
RESULT 5
Q8LVE1
                                   PRT;
                                           16 AA.
ΙD
    Q8LVE1
                 PRELIMINARY;
AC
     Q8LVE1;
DT
     01-OCT-2002 (TrEMBLrel. 22, Created)
     01-OCT-2002 (TrEMBLrel. 22, Last sequence update)
DT
     01-JUN-2003 (TrEMBLrel. 24, Last annotation update)
DT
DΕ
     PsaA (Fragment).
GN
     PSAA.
os
    Punica granatum (Pomegranate).
OG
     Chloroplast.
OC
     Eukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta;
OC
     Spermatophyta; Magnoliophyta; eudicotyledons; core eudicots; rosids;
    Myrtales; Lythraceae; Punica.
OC
OX
    NCBI TaxID=22663;
RN
     [1]
RP
     SEQUENCE FROM N.A.
RC
     STRAIN=P728, and P745;
RA
     Huang Y., Shi S.;
RT
     "Phylogenetics of Lythraceae sensu lato: A Preliminary Analysis Based
     on Chloroplast rbcL Gene, psaA-ycf3 Spacer, and Nuclear rDNA Internal
RT
     Transcribed Spacer (ITS) Sequences.";
RT
     Int. J. Plant Sci. 163:215-225(2002).
RL
     EMBL; AY035724; AAL14129.1; -.
DR
     EMBL; AY035742; AAL14165.1; -.
DR
     GO; GO:0009507; C:chloroplast; IEA.
DR
KW
     Chloroplast.
FT
     NON TER
                  16
                         16
     SEQUENCE
               16 AA; 1895 MW;
                                 517FE691B89355B9 CRC64;
SQ
  Query Match
                          30.3%; Score 27; DB 8; Length 16;
  Best Local Similarity 15.4%; Pred. No. 1.4e+03;
  Matches
             2; Conservative 10; Mismatches 1; Indels
                                                                 0; Gaps
                                                                             0;
            1 LQTPQPLLQVMME 13
Qу
              :::|:| ::::::
Db
            3 IRSPEPEVKILVD 15
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RESULT 6
08MC53
ID
     Q8MC53
                 PRELIMINARY;
                                    PRT;
                                            16 AA.
AC
     08MC53;
DT
     01-OCT-2002 (TrEMBLrel. 22, Created)
     01-OCT-2002 (TrEMBLrel. 22, Last sequence update)
DT
     01-JUN-2003 (TrEMBLrel. 24, Last annotation update)
DT
DE
     PsaA (Fragment).
GN
     PSAA.
     Woodfordia fruticosa.
os
OG
     Chloroplast.
     Eukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta;
OC
     Spermatophyta; Magnoliophyta; eudicotyledons; core eudicots; rosids;
OC
OC
     Myrtales; Lythraceae; Woodfordia.
OX
     NCBI TaxID=141189;
RN
     [1]
RP
     SEQUENCE FROM N.A.
     STRAIN=W493;
RC
RA
     Huang Y., Shi S.;
     "Phylogenetics of Lythraceae sensu lato: A Preliminary Analysis Based
RT
RT
     on Chloroplast rbcL Gene, psaA-ycf3 Spacer, and Nuclear rDNA Internal
RT
     Transcribed Spacer (ITS) Sequences.";
     Int. J. Plant Sci. 163:215-225(2002).
RL
DR
     EMBL; AY035722; AAL14125.1; -.
     GO; GO:0009507; C:chloroplast; IEA.
DR
KW
     Chloroplast.
     NON TER
FT
                  16
                         16
     SEQUENCE
                        1895 MW; 517FE691B89355B9 CRC64;
SQ
                16 AA;
                          30.3%; Score 27; DB 8; Length 16;
  Query Match
  Best Local Similarity
                                   Pred. No. 1.4e+03;
                          15.4%;
  Matches
             2; Conservative
                               10; Mismatches
                                                    1; Indels
                                                                   0; Gaps
                                                                               0;
            1 LQTPQPLLQVMME 13
Qу
              :::|:| ::::::
Db
            3 IRSPEPEVKILVD 15
RESULT 7
Q8MC17
ID
     Q8MC17
                 PRELIMINARY;
                                    PRT;
                                            16 AA.
AC
     Q8MC17;
DT
     01-OCT-2002 (TrEMBLrel. 22, Created)
     01-OCT-2002 (TrEMBLrel. 22, Last sequence update)
DT
     01-JUN-2003 (TrEMBLrel. 24, Last annotation update)
\mathbf{DT}
DE
     PsaA (Fragment).
     PSAA.
GN
os
     Ludwigia hyssopifolia.
OG
     Chloroplast.
     Eukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta;
OC
     Spermatophyta; Magnoliophyta; eudicotyledons; core eudicots; rosids;
OC
     Myrtales; Onagraceae; Ludwigia.
OC
OX
     NCBI TaxID=155013;
RN
     [1]
RP
     SEQUENCE FROM N.A.
RC
     STRAIN=L787;
     Huang Y., Shi S.;
RA
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RT
     "Phylogenetics of Lythraceae sensu lato: A Preliminary Analysis Based
     on Chloroplast rbcL Gene, psaA-ycf3 Spacer, and Nuclear rDNA Internal
RT
RT
     Transcribed Spacer (ITS) Sequences.";
     Int. J. Plant Sci. 163:215-225(2002).
RL
     EMBL; AY035745; AAL14171.1; -.
DR
DR
     GO; GO:0009507; C:chloroplast; IEA.
KW
     Chloroplast.
FT
     NON TER
                  16
                         16
     SEQUENCE
SO
                16 AA; 1895 MW; 517FE691B89355B9 CRC64;
                          30.3%; Score 27; DB 8; Length 16;
  Query Match
                          15.4%; Pred. No. 1.4e+03;
  Best Local Similarity
             2; Conservative 10; Mismatches 1; Indels
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                                                                              0;
Qу
            1 LQTPQPLLQVMME 13
              :::|:| ::::::
Db
            3 IRSPEPEVKILVD 15
RESULT 8
Q8LVE2
                                           16 AA.
                 PRELIMINARY;
                                   PRT:
ID
    Q8LVE2
AC
     O8LVE2;
     01-OCT-2002 (TrEMBLrel. 22, Created)
DT
     01-OCT-2002 (TrEMBLrel. 22, Last sequence update)
DT
     01-JUN-2003 (TrEMBLrel. 24, Last annotation update)
DT
DE
     PsaA (Fragment).
     PSAA.
GN
     Lythrum salicaria (Purple loosestrife).
OS
OG
     Chloroplast.
     Eukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta;
OC
     Spermatophyta; Magnoliophyta; eudicotyledons; core eudicots; rosids;
OC
     Myrtales; Lythraceae; Lythrum.
OC
OX
     NCBI TaxID=13129;
RN
     [1]
     SEQUENCE FROM N.A.
RP
     STRAIN=L758, and 758F;
RC
     Huang Y., Shi S.;
RA
RT
     "Phylogenetics of Lythraceae sensu lato: A Preliminary Analysis Based
RT
     on Chloroplast rbcL Gene, psaA-ycf3 Spacer, and Nuclear rDNA Internal
RT
     Transcribed Spacer (ITS) Sequences.";
     Int. J. Plant Sci. 163:215-225(2002).
RL
DR
     EMBL; AY035727; AAL14135.1; -.
     EMBL; AF421495; AAM45853.1; -.
DR
     GO; GO:0009507; C:chloroplast; IEA.
DR
KW
     Chloroplast.
FT
     NON TER
                  16
                         16
SQ
     SEQUENCE
                16 AA; 1895 MW; 517FE691B89355B9 CRC64;
                                  Score 27; DB 8; Length 16;
  Query Match
                          30.3%;
                                  Pred. No. 1.4e+03;
  Best Local Similarity 15.4%;
  Matches
             2; Conservative 10; Mismatches
                                                   1; Indels
                                                                  0; Gaps
                                                                              0;
Qу
            1 LOTPOPLLOVMME 13
              :::|:| ::::::
Db
            3 IRSPEPEVKILVD 15
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RESULT 9
Q8LVE0
ID
                 PRELIMINARY;
                                   PRT:
                                           16 AA.
    Q8LVE0
AC
    Q8LVE0;
     01-OCT-2002 (TrEMBLrel. 22, Created)
DT
     01-OCT-2002 (TrEMBLrel. 22, Last sequence update)
DΤ
    01-JUN-2003 (TrEMBLrel. 24, Last annotation update)
DT
DE
    PsaA (Fragment).
GN
    PSAA.
    Trapa maximowiczii.
OS
OG
    Chloroplast.
    Eukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta;
OC
    Spermatophyta; Magnoliophyta; eudicotyledons; core eudicots; rosids;
OC
OC
    Myrtales; Lythraceae; Trapa.
OX
    NCBI TaxID=162053;
RN
     [1]
RP
     SEQUENCE FROM N.A.
RC
    STRAIN=T744, and T010;
    Huang Y., Shi S.;
RA
     "Phylogenetics of Lythraceae sensu lato: A Preliminary Analysis Based
RT
    on Chloroplast rbcL Gene, psaA-ycf3 Spacer, and Nuclear rDNA Internal
RT
    Transcribed Spacer (ITS) Sequences.";
RT
    Int. J. Plant Sci. 163:215-225(2002).
RL
    EMBL; AY035729; AAL14139.1; -.
DR
    EMBL; AY035730; AAL14141.1; -.
DR
    GO; GO:0009507; C:chloroplast; IEA.
DR
KW
    Chloroplast.
FT
    NON TER
                  16
                         16
     SEQUENCE
                16 AA; 1895 MW; 517FE691B89355B9 CRC64;
SO
                          30.3%; Score 27; DB 8; Length 16;
  Query Match
  Best Local Similarity 15.4%; Pred. No. 1.4e+03;
  Matches
             2; Conservative 10; Mismatches 1; Indels
                                                                  0; Gaps
                                                                              0;
            1 LQTPQPLLQVMME 13
Qу
              :::|:| ::::::
Dh
            3 IRSPEPEVKILVD 15
RESULT 10
Q8MET2
ID
     Q8MET2
                 PRELIMINARY;
                                   PRT;
                                           16 AA.
AC
     Q8MET2;
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DT
     01-OCT-2002 (TrEMBLrel. 22, Last sequence update)
DT
DT
     01-JUN-2003 (TrEMBLrel. 24, Last annotation update)
DE
     PSI P700 apoprotein Al (Fragment).
GN
     PSAA.
OS
     Saxifraga stolonifera.
OG
     Chloroplast.
OC
     Eukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta;
     Spermatophyta; Magnoliophyta; eudicotyledons; core eudicots;
OC
OC
     Saxifragales; Saxifragaceae; Saxifraga.
OX
     NCBI TaxID=182070;
RN
     [1]
     SEQUENCE FROM N.A.
RP
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RA
     Shi S., Huang Y., Zhong Y., Du Y., Zhang Q., Chang H., Boufford D.E.;
RT
     "Phylogeny of the Altingiaceae based on cpDNA matK, PY-IGS and nrDNA
     ITS sequences.";
RT
     Plant Syst. Evol. 230:13-24(2001).
RL
     EMBL; AF377995; AAM45512.1; -.
DR
DR
     GO; GO:0009507; C:chloroplast; IEA.
     Chloroplast.
KW
FT
     NON TER
                  16
                         16
     SEQUENCE
                16 AA; 1895 MW; 517FE691B89355B9 CRC64;
SQ
                          30.3%; Score 27; DB 8; Length 16;
  Query Match
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  Best Local Similarity
             2; Conservative 10; Mismatches
  Matches
                                                   1; Indels
                                                                 0; Gaps
                                                                              0;
            1 LQTPQPLLQVMME 13
Qу
              :::|:| ::::::
Db
            3 IRSPEPEVKILVD 15
RESULT 11
08MC45
ID
                                           16 AA.
    Q8MC45
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AC
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DT
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     01-OCT-2002 (TrEMBLrel. 22, Last sequence update)
DT
     01-JUN-2003 (TrEMBLrel. 24, Last annotation update)
DT
DΕ
    PsaA (Fragment).
GN
    PSAA.
OS
    Decodon verticillatus (Swamp loosestrife).
OG
    Chloroplast.
OC
     Eukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta;
OC
     Spermatophyta; Magnoliophyta; eudicotyledons; core eudicots; rosids;
OC
    Myrtales; Lythraceae; Decodon.
OX
    NCBI TaxID=162018;
RN
     [1]
RP
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RC
     STRAIN=D212;
RA
    Huang Y., Shi S.;
RT
     "Phylogenetics of Lythraceae sensu lato: A Preliminary Analysis Based
RT
     on Chloroplast rbcL Gene, psaA-ycf3 Spacer, and Nuclear rDNA Internal
RT
     Transcribed Spacer (ITS) Sequences.";
RL
    Int. J. Plant Sci. 163:215-225(2002).
    EMBL; AY035728; AAL14137.1; -.
DR
DR
    GO; GO:0009507; C:chloroplast; IEA.
KW
     Chloroplast.
FT
     NON TER
                  16
                         16
     SEQUENCE
SQ
                16 AA; 1895 MW;
                                  517FE691B89355B9 CRC64;
  Query Match
                          30.3%;
                                  Score 27; DB 8; Length 16;
  Best Local Similarity 15.4%; Pred. No. 1.4e+03;
 Matches
             2; Conservative
                              10; Mismatches
                                                1; Indels
                                                                 0; Gaps
                                                                              0;
            1 LQTPQPLLQVMME 13
Qy
             :::|:| ::::::
Db
            3 IRSPEPEVKILVD 15
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RESULT 12
Q8MC21
ID
     Q8MC21
                 PRELIMINARY;
                                   PRT:
                                            16 AA.
AC
     Q8MC21;
     01-OCT-2002 (TrEMBLrel. 22, Created)
DT
     01-OCT-2002 (TrEMBLrel. 22, Last sequence update)
DT
DT
     01-JUN-2003 (TrEMBLrel. 24, Last annotation update)
DE
     PsaA (Fragment).
GN
     PSAA.
os
     Combretum wallichii.
OG
     Chloroplast.
     Eukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta;
OC
OC
     Spermatophyta; Magnoliophyta; eudicotyledons; core eudicots; rosids;
OC
     Myrtales; Combretaceae; Combretum.
     NCBI TaxID=131243;
OX
RN
    [1]
RP
     SEOUENCE FROM N.A.
RC
     STRAIN=C505;
RA
     Huang Y., Shi S.;
     "Phylogenetics of Lythraceae sensu lato: A Preliminary Analysis Based
RT
     on Chloroplast rbcL Gene, psaA-ycf3 Spacer, and Nuclear rDNA Internal
RT
RT
     Transcribed Spacer (ITS) Sequences.";
     Int. J. Plant Sci. 163:215-225(2002).
RL
DR
     EMBL; AY035743; AAL14167.1; -.
     GO; GO:0009507; C:chloroplast; IEA.
DR
KW
     Chloroplast.
     NON TER
                  16
                         16
FT
SQ
     SEQUENCE
                16 AA; 1895 MW; 517FE691B89355B9 CRC64;
                          30.3%; Score 27; DB 8; Length 16;
  Query Match
  Best Local Similarity 15.4%; Pred. No. 1.4e+03;
             2; Conservative
                                10; Mismatches
                                                  1; Indels
                                                                  0;
                                                                      Gaps
                                                                               0;
            1 LQTPQPLLQVMME 13
Qy
              :::|:| ::::::
Db
            3 IRSPEPEVKILVD 15
RESULT 13
Q8MC51
     Q8MC51
ΙD
                 PRELIMINARY;
                                   PRT:
                                            16 AA.
AC
     Q8MC51;
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DT
     01-OCT-2002 (TrEMBLrel. 22, Last sequence update)
DT
DT
     01-JUN-2003 (TrEMBLrel. 24, Last annotation update)
DE
     PsaA (Fragment).
GN
     PSAA.
os
     Cuphea lanceolata.
OG
     Chloroplast.
     Eukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta;
OC
OC
     Spermatophyta; Magnoliophyta; eudicotyledons; core eudicots; rosids;
OC
     Myrtales; Lythraceae; Cuphea.
OX
     NCBI TaxID=3930;
RN
     [1]
RP
     SEQUENCE FROM N.A.
RC
     STRAIN=C490;
RA
     Huang Y., Shi S.;
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"Phylogenetics of Lythraceae sensu lato: A Preliminary Analysis Based
RT
RT
     on Chloroplast rbcL Gene, psaA-ycf3 Spacer, and Nuclear rDNA Internal
RT
     Transcribed Spacer (ITS) Sequences.";
RL
     Int. J. Plant Sci. 163:215-225(2002).
DR
     EMBL; AY035723; AAL14127.1; -.
DR
     GO; GO:0009507; C:chloroplast; IEA.
KW
     Chloroplast.
     NON TER
FT
                  16
                         16
     SEQUENCE
                16 AA; 1895 MW; 517FE691B89355B9 CRC64;
SQ
  Query Match
                          30.3%; Score 27; DB 8; Length 16;
                         15.4%; Pred. No. 1.4e+03;
  Best Local Similarity
             2; Conservative 10; Mismatches
  Matches
                                                  1; Indels
                                                                 0; Gaps
                                                                             0;
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Qу
              :::|:| ::::::
Db
            3 IRSPEPEVKILVD 15
RESULT 14
Q8MC33
ID
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                                           16 AA.
AC
     Q8MC33;
DΤ
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DT
     01-OCT-2002 (TrEMBLrel. 22, Last sequence update)
DT
     01-JUN-2003 (TrEMBLrel. 24, Last annotation update)
DΕ
     PsaA (Fragment).
GN
     PSAA.
os
     Rotala indica.
OG
     Chloroplast.
OC
     Eukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta;
OC
     Spermatophyta; Magnoliophyta; eudicotyledons; core eudicots; rosids;
OC
     Myrtales; Lythraceae; Rotala.
OX
     NCBI TaxID=162024;
RN
     [1]
     SEQUENCE FROM N.A.
RP
RC
     STRAIN=R492;
RA
     Huang Y., Shi S.;
RT
     "Phylogenetics of Lythraceae sensu lato: A Preliminary Analysis Based
RT
     on Chloroplast rbcL Gene, psaA-ycf3 Spacer, and Nuclear rDNA Internal
     Transcribed Spacer (ITS) Sequences.";
RT
RL
     Int. J. Plant Sci. 163:215-225(2002).
DR
     EMBL; AY035736; AAL14153.1; -.
DR
     GO; GO:0009507; C:chloroplast; IEA.
KW
     Chloroplast.
FT
     NON TER
                  16
                         16
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SQ
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                                  Score 27; DB 8; Length 16;
  Best Local Similarity 15.4%; Pred. No. 1.4e+03;
                              10; Mismatches 1; Indels
  Matches
             2; Conservative
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                                                                             0;
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Qу
             :::|:| ::::::
Db
            3 IRSPEPEVKILVD 15
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RESULT 15
Q8MC19
ID
     Q8MC19
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AC
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DT
     01-OCT-2002 (TrEMBLrel. 22, Created)
     01-OCT-2002 (TrEMBLrel. 22, Last sequence update)
DT
     01-JUN-2003 (TrEMBLrel. 24, Last annotation update)
DT
DE
     PsaA (Fragment).
     PSAA.
GN
os
     Quisqualis indica (Rangoon creeper).
OG
     Chloroplast.
OC
     Eukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta;
     Spermatophyta; Magnoliophyta; eudicotyledons; core eudicots; rosids;
OC
OC
     Myrtales; Combretaceae; Quisqualis.
     NCBI TaxID=3956;
OX
RN
     [1]
RP
     SEQUENCE FROM N.A.
RC
     STRAIN=Q379;
RA
     Huang Y., Shi S.;
     "Phylogenetics of Lythraceae sensu lato: A Preliminary Analysis Based
RT
RT
     on Chloroplast rbcL Gene, psaA-ycf3 Spacer, and Nuclear rDNA Internal
RT
     Transcribed Spacer (ITS) Sequences.";
RL
     Int. J. Plant Sci. 163:215-225(2002).
DR
     EMBL; AY035744; AAL14169.1; -.
     GO; GO:0009507; C:chloroplast; IEA.
DR
KW
     Chloroplast.
FT
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                  16
                         16
     SEQUENCE
SO
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  Query Match
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  Best Local Similarity
                          15.4%; Pred. No. 1.4e+03;
             2; Conservative
                               10: Mismatches
                                                  1; Indels
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                                                                      Gaps
                                                                              0:
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Qу
              :::|:| ::::::
Db
            3 IRSPEPEVKILVD 15
RESULT 16
Q8MC49
ID
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                 PRELIMINARY;
                                   PRT:
                                            16 AA.
AC
     Q8MC49;
DT
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     01-OCT-2002 (TrEMBLrel. 22, Last sequence update)
DT
     01-JUN-2003 (TrEMBLrel. 24, Last annotation update)
DT
DE
     PsaA (Fragment).
GN
     PSAA.
os
     Pemphis acidula.
OG
     Chloroplast.
OC
     Eukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta;
OC
     Spermatophyta; Magnoliophyta; eudicotyledons; core eudicots; rosids;
OC
     Myrtales; Lythraceae; Pemphis.
     NCBI TaxID=126635;
OX
RN
     [1]
RP
     SEQUENCE FROM N.A.
RC
     STRAIN=P630;
RA
     Huang Y., Shi S.;
```

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RT
     "Phylogenetics of Lythraceae sensu lato: A Preliminary Analysis Based
RT
     on Chloroplast rbcL Gene, psaA-ycf3 Spacer, and Nuclear rDNA Internal
RT
    Transcribed Spacer (ITS) Sequences.";
RL
     Int. J. Plant Sci. 163:215-225(2002).
DR
     EMBL; AY035725; AAL14131.1; -.
DR
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KW
     Chloroplast.
    NON TER
FT
                  16
                        16
SQ
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  Query Match
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  Best Local Similarity 15.4%; Pred. No. 1.4e+03;
            2; Conservative 10; Mismatches
 Matches
                                                 1; Indels
                                                                 0; Gaps
                                                                             0;
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Qу
             :::|:| ::::::
Db
            3 IRSPEPEVKILVD 15
RESULT 17
O8MC27
ID
                                   PRT:
                                           16 AA.
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AC
    Q8MC27;
DT
     01-OCT-2002 (TrEMBLrel. 22, Created)
     01-OCT-2002 (TrEMBLrel. 22, Last sequence update)
     01-JUN-2003 (TrEMBLrel. 24, Last annotation update)
DT
DE
    PsaA (Fragment).
GN
    PSAA.
OS
    Lagerstroemia villosa.
OG
    Chloroplast.
OC
     Eukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta;
OC
     Spermatophyta; Magnoliophyta; eudicotyledons; core eudicots; rosids;
OC
    Myrtales; Lythraceae; Lagerstroemia.
OX
    NCBI TaxID=162025;
RN
     [1]
RP
     SEQUENCE FROM N.A.
RC
     STRAIN=L688;
    Huang Y., Shi S.;
RA
RT
     "Phylogenetics of Lythraceae sensu lato: A Preliminary Analysis Based
RT
     on Chloroplast rbcL Gene, psaA-ycf3 Spacer, and Nuclear rDNA Internal
    Transcribed Spacer (ITS) Sequences.";
RT
    Int. J. Plant Sci. 163:215-225(2002).
RL
    EMBL; AY035739; AAL14159.1; -.
DR
DR
    GO; GO:0009507; C:chloroplast; IEA.
KW
    Chloroplast.
    NON TER
FT
                  16
                        16
     SEQUENCE
SQ
               16 AA; 1895 MW; 517FE691B89355B9 CRC64;
  Query Match
                          30.3%; Score 27; DB 8; Length 16;
  Best Local Similarity 15.4%; Pred. No. 1.4e+03;
  Matches
             2; Conservative
                              10; Mismatches
                                                1; Indels
                                                                 0; Gaps
                                                                             0;
            1 LQTPQPLLQVMME 13
Qу
             :::|:| ::::::
           3 IRSPEPEVKILVD 15
Db
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RESULT 18
O8METO
ID
     Q8MET0
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AC
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     01-OCT-2002 (TrEMBLrel. 22, Created)
DT
     01-OCT-2002 (TrEMBLrel. 22, Last sequence update)
DT
     01-JUN-2003 (TrEMBLrel. 24, Last annotation update)
DT
DE
     PSI P700 apoprotein A1 (Fragment).
GN
     PSAA.
     Cercidiphyllum japonicum (Katsura tree).
OS
OG
     Chloroplast.
     Eukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta;
OC
OC
     Spermatophyta; Magnoliophyta; eudicotyledons; core eudicots;
OC
     Saxifragales; Cercidiphyllaceae; Cercidiphyllum.
OX
     NCBI TaxID=13413;
RN
     [1]
RP
     SEQUENCE FROM N.A.
RA
     Shi S., Huang Y., Zhong Y., Du Y., Zhang Q., Chang H., Boufford D.E.;
RT
     "Phylogeny of the Altingiaceae based on cpDNA matk, PY-IGS and nrDNA
RT
     ITS sequences.";
     Plant Syst. Evol. 230:13-24(2001).
RL
DR
     EMBL; AF377996; AAM45514.1; -.
DR
     GO; GO:0009507; C:chloroplast; IEA.
KW
     Chloroplast.
FT
    NON TER
                  16
                         16
     SEQUENCE
                16 AA; 1895 MW; 517FE691B89355B9 CRC64;
SQ
  Query Match
                          30.3%; Score 27; DB 8; Length 16;
 Best Local Similarity
                          15.4%; Pred. No. 1.4e+03;
 Matches
                               10; Mismatches 1; Indels
            2; Conservative
            1 LQTPQPLLQVMME 13
Qу
              :::|:| ::::::
            3 IRSPEPEVKILVD 15
Db
RESULT 19
Q8MC23
ID
    Q8MC23
                 PRELIMINARY;
                                   PRT;
                                           16 AA.
AC
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DT
     01-OCT-2002 (TrEMBLrel. 22, Created)
DΤ
     01-OCT-2002 (TrEMBLrel. 22, Last sequence update)
     01-JUN-2003 (TrEMBLrel. 24, Last annotation update)
DT
DE
    PsaA (Fragment).
    PSAA.
GN
OS
     Sonneratia alba.
OG
     Chloroplast.
OC
     Eukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta;
OC
     Spermatophyta; Magnoliophyta; eudicotyledons; core eudicots; rosids;
OC
    Myrtales; Lythraceae; Sonneratia.
OX
    NCBI TaxID=122812;
RN
     [1]
RP
     SEQUENCE FROM N.A.
RC
     STRAIN=S482;
RA
     Huang Y., Shi S.;
RT
     "Phylogenetics of Lythraceae sensu lato: A Preliminary Analysis Based
RT
     on Chloroplast rbcL Gene, psaA-ycf3 Spacer, and Nuclear rDNA Internal
```

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Transcribed Spacer (ITS) Sequences.";
RT
     Int. J. Plant Sci. 163:215-225(2002).
RL
     EMBL; AY035741; AAL14163.1; -.
DR
     GO; GO:0009507; C:chloroplast; IEA.
DR
KW
     Chloroplast.
FT
     NON TER
                  16
                         16
     SEQUENCE
SO
                16 AA; 1895 MW; 517FE691B89355B9 CRC64;
  Query Match
                          30.3%; Score 27; DB 8; Length 16;
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            2; Conservative 10; Mismatches
                                                                             0;
                                                  1; Indels
                                                                 0; Gaps
  Matches
            1 LQTPQPLLQVMME 13
Qу
              :::|:| ::::::
            3 IRSPEPEVKILVD 15
Db
RESULT 20
Q8MC41
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                                           16 AA.
ID
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                 PRELIMINARY;
AC
     Q8MC41;
     01-OCT-2002 (TrEMBLrel. 22, Created)
DT
     01-OCT-2002 (TrEMBLrel. 22, Last sequence update)
     01-JUN-2003 (TrEMBLrel. 24, Last annotation update)
DT
DE
     PsaA (Fragment).
GN
     PSAA.
     Nesaea luederitzii.
os
OG
     Chloroplast.
     Eukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta;
OC
     Spermatophyta; Magnoliophyta; eudicotyledons; core eudicots; rosids;
OC
OC
     Myrtales; Lythraceae; Nesaea.
OX
     NCBI TaxID=162020;
RN
     [1]
RP
     SEQUENCE FROM N.A.
RC
     STRAIN=N213;
RA
     Huang Y., Shi S.;
     "Phylogenetics of Lythraceae sensu lato: A Preliminary Analysis Based
RT
     on Chloroplast rbcL Gene, psaA-ycf3 Spacer, and Nuclear rDNA Internal
RT
RT
     Transcribed Spacer (ITS) Sequences.";
RL
     Int. J. Plant Sci. 163:215-225(2002).
DR
     EMBL; AY035732; AAL14145.1; -.
     GO; GO:0009507; C:chloroplast; IEA.
DR
KW
     Chloroplast.
     NON TER
FT
                  16
                         16
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SQ
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  Best Local Similarity 15.4%; Pred. No. 1.4e+03;
                              10; Mismatches 1; Indels
                                                                             0;
  Matches
             2; Conservative
                                                                 0; Gaps
            1 LOTPOPLLOVMME 13
Qу
              :::|:| ::::::
Db
            3 IRSPEPEVKILVD 15
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RESULT 21 Q8MC15

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ΙD
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                                   PRT:
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AC
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DT
     01-OCT-2002 (TrEMBLrel. 22, Created)
     01-OCT-2002 (TrEMBLrel. 22, Last sequence update)
DT
     01-JUN-2003 (TrEMBLrel. 24, Last annotation update)
DΤ
DE
     PsaA (Fragment).
    PSAA.
GN
    Fuchsia hybrid cultivar.
os
OG
    Chloroplast.
     Eukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta;
OC
OC
     Spermatophyta; Magnoliophyta; eudicotyledons; core eudicots; rosids;
    Myrtales; Onagraceae; Fuchsia.
OC
OX
    NCBI TaxID=133545;
RN
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RP
    SEQUENCE FROM N.A.
RC
    STRAIN=F016;
RA
    Huang Y., Shi S.;
RT
     "Phylogenetics of Lythraceae sensu lato: A Preliminary Analysis Based
     on Chloroplast rbcL Gene, psaA-ycf3 Spacer, and Nuclear rDNA Internal
RT
    Transcribed Spacer (ITS) Sequences.";
RT
    Int. J. Plant Sci. 163:215-225(2002).
RL
    EMBL; AY035746; AAL14173.1; -.
DR
DR
    GO; GO:0009507; C:chloroplast; IEA.
KW
    Chloroplast.
FT
    NON TER
                  16
                         16
     SEQUENCE
                16 AA; 1895 MW; 517FE691B89355B9 CRC64;
SQ
 Query Match
                          30.3%; Score 27; DB 8; Length 16;
  Best Local Similarity
                          15.4%; Pred. No. 1.4e+03;
           2; Conservative 10; Mismatches 1; Indels
 Matches
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                                                                              0;
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Qу
              :::|:| ::::::
            3 IRSPEPEVKILVD 15
Db
RESULT 22
Q8MC43
ID
    Q8MC43
                 PRELIMINARY;
                                   PRT;
                                           16 AA.
AC
    O8MC43;
DT
     01-OCT-2002 (TrEMBLrel. 22, Created)
     01-OCT-2002 (TrEMBLrel. 22, Last sequence update)
DT
     01-JUN-2003 (TrEMBLrel. 24, Last annotation update)
DT
DE
    PsaA (Fragment).
    PSAA.
GN
os
     Sonneratia caseolaris.
OG
     Chloroplast.
OC
    Eukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta;
OC
     Spermatophyta; Magnoliophyta; eudicotyledons; core eudicots; rosids;
    Myrtales; Lythraceae; Sonneratia.
OC
OX
    NCBI TaxID=122814;
RN
     [1]
RP
    SEQUENCE FROM N.A.
RC
     STRAIN=S435;
RA
     Huang Y., Shi S.;
RT
     "Phylogenetics of Lythraceae sensu lato: A Preliminary Analysis Based
RT
     on Chloroplast rbcL Gene, psaA-ycf3 Spacer, and Nuclear rDNA Internal
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RT
     Transcribed Spacer (ITS) Sequences.";
     Int. J. Plant Sci. 163:215-225(2002).
RL
DR
     EMBL; AY035731; AAL14143.1; -.
DR
     GO; GO:0009507; C:chloroplast; IEA.
KW
     Chloroplast.
     NON TER
FT
                  16
                         16
     SEQUENCE
                16 AA; 1895 MW; 517FE691B89355B9 CRC64;
SO
  Ouery Match
                          30.3%;
                                  Score 27; DB 8; Length 16;
                          15.4%; Pred. No. 1.4e+03;
  Best Local Similarity
             2; Conservative 10; Mismatches
  Matches
                                                  1; Indels
                                                                              0;
                                                                 0; Gaps
            1 LQTPQPLLQVMME 13
Qу
              :::|:| ::::::
            3 IRSPEPEVKILVD 15
Db
RESULT 23
Q8MC47
ΙD
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                 PRELIMINARY;
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                                           16 AA.
AC
     Q8MC47;
DT
     01-OCT-2002 (TrEMBLrel. 22, Created)
DT
     01-OCT-2002 (TrEMBLrel. 22, Last sequence update)
     01-JUN-2003 (TrEMBLrel. 24, Last annotation update)
DE
     PsaA (Fragment).
GN
     PSAA.
     Peplis portula.
OS
OG
     Chloroplast.
OC
     Eukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta;
OC
     Spermatophyta; Magnoliophyta; eudicotyledons; core eudicots; rosids;
     Myrtales; Lythraceae; Peplis.
OC
OX
     NCBI TaxID=162016;
RN
     [1]
RP
     SEQUENCE FROM N.A.
RC
     STRAIN=P220;
RA
     Huang Y., Shi S.;
RT
     "Phylogenetics of Lythraceae sensu lato: A Preliminary Analysis Based
RT
     on Chloroplast rbcL Gene, psaA-ycf3 Spacer, and Nuclear rDNA Internal
RT
     Transcribed Spacer (ITS) Sequences.";
RL
     Int. J. Plant Sci. 163:215-225(2002).
DR
     EMBL; AY035726; AAL14133.1; -.
     GO; GO:0009507; C:chloroplast; IEA.
DR
KW
     Chloroplast.
     NON TER
FT
                  16
                         16
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                16 AA; 1895 MW;
SQ
                                 517FE691B89355B9 CRC64;
  Query Match
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                                  Score 27; DB 8; Length 16;
  Best Local Similarity 15.4%; Pred. No. 1.4e+03;
  Matches
             2; Conservative
                              10; Mismatches
                                                 1; Indels
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Qу
              :::|:| ::::::
Db
            3 IRSPEPEVKILVD 15
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RESULT 24 Q8MC37

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08MC37
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                                           16 AA.
ID
                 PRELIMINARY:
AC
    Q8MC37;
DT
     01-OCT-2002 (TrEMBLrel. 22, Created)
     01-OCT-2002 (TrEMBLrel. 22, Last sequence update)
     01-JUN-2003 (TrEMBLrel. 24, Last annotation update)
DT
DΕ
     PsaA (Fragment).
GN
     PSAA.
OS
    Lawsonia inermis.
    Chloroplast.
OG
OC
     Eukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta;
     Spermatophyta; Magnoliophyta; eudicotyledons; core eudicots; rosids;
OC
    Myrtales; Lythraceae; Lawsonia.
OC
    NCBI TaxID=141191;
OX
RN
     [1]
    SEQUENCE FROM N.A.
RP
RC
    STRAIN=L494;
RA
    Huang Y., Shi S.;
RT
     "Phylogenetics of Lythraceae sensu lato: A Preliminary Analysis Based
RT
     on Chloroplast rbcL Gene, psaA-ycf3 Spacer, and Nuclear rDNA Internal
     Transcribed Spacer (ITS) Sequences.";
RT
RL
     Int. J. Plant Sci. 163:215-225(2002).
DR
    EMBL; AY035734; AAL14149.1; -.
    GO; GO:0009507; C:chloroplast; IEA.
DR
KW
    Chloroplast.
    NON TER
                  16
                         16
FT
    SEQUENCE
               16 AA; 1895 MW; 517FE691B89355B9 CRC64;
SQ
                          30.3%; Score 27; DB 8; Length 16;
 Query Match
  Best Local Similarity
                          15.4%; Pred. No. 1.4e+03;
            2; Conservative 10; Mismatches 1; Indels
  Matches
                                                                 0; Gaps
                                                                              0;
Qу
            1 LOTPOPLLOVMME 13
             :::|:| ::::::
Db
            3 IRSPEPEVKILVD 15
RESULT 25
08MC29
ID
    Q8MC29
                 PRELIMINARY;
                                   PRT;
                                           16 AA.
AC
     01-OCT-2002 (TrEMBLrel. 22, Created)
DT
     01-OCT-2002 (TrEMBLrel. 22, Last sequence update)
    01-JUN-2003 (TrEMBLrel. 24, Last annotation update)
DT
DE
    PsaA (Fragment).
GN
    PSAA.
os
    Duabanga grandiflora.
OG
    Chloroplast.
OC
     Eukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta;
OC
     Spermatophyta; Magnoliophyta; eudicotyledons; core eudicots; rosids;
OC
    Myrtales; Lythraceae; Duabanga.
OX
    NCBI TaxID=122808;
RN
     [1]
RP
    SEQUENCE FROM N.A.
RC
    STRAIN=D441;
RA
     Huang Y., Shi S.;
RT
     "Phylogenetics of Lythraceae sensu lato: A Preliminary Analysis Based
RT
     on Chloroplast rbcL Gene, psaA-ycf3 Spacer, and Nuclear rDNA Internal
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Transcribed Spacer (ITS) Sequences.";
RT
     Int. J. Plant Sci. 163:215-225(2002).
RL
DR
     EMBL; AY035738; AAL14157.1; -.
     GO; GO:0009507; C:chloroplast; IEA.
DR
KW
     Chloroplast.
     NON TER
\mathbf{FT}
                  16
                         16
SQ
     SEQUENCE
                16 AA; 1895 MW; 517FE691B89355B9 CRC64;
                          30.3%; Score 27; DB 8; Length 16;
  Query Match
  Best Local Similarity 15.4%; Pred. No. 1.4e+03;
             2; Conservative 10; Mismatches
                                                  1; Indels
                                                                  0; Gaps
                                                                              0;
            1 LOTPOPLLOVMME 13
Qу
              :::|:| ::::::
Db
            3 IRSPEPEVKILVD 15
RESULT 26
Q8MC31
ID
     Q8MC31
                 PRELIMINARY;
                                   PRT;
                                           16 AA.
     Q8MC31;
AC
DT
     01-OCT-2002 (TrEMBLrel. 22, Created)
     01-OCT-2002 (TrEMBLrel. 22, Last sequence update)
DT
DT
     01-JUN-2003 (TrEMBLrel. 24, Last annotation update)
DE
     PsaA (Fragment).
    PSAA.
GN
os
     Lagerstroemia speciosa.
OG
     Chloroplast.
     Eukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta;
OC
     Spermatophyta; Magnoliophyta; eudicotyledons; core eudicots; rosids;
OC
OC
     Myrtales; Lythraceae; Lagerstroemia.
OX
     NCBI TaxID=122810;
RN
     [1]
RP
     SEQUENCE FROM N.A.
     STRAIN=L477;
RC
RA
     Huang Y., Shi S.;
     "Phylogenetics of Lythraceae sensu lato: A Preliminary Analysis Based
RT
     on Chloroplast rbcL Gene, psaA-ycf3 Spacer, and Nuclear rDNA Internal
RT
RT
     Transcribed Spacer (ITS) Sequences.";
RL
     Int. J. Plant Sci. 163:215-225(2002).
DR
     EMBL; AY035737; AAL14155.1; -.
     GO; GO:0009507; C:chloroplast; IEA.
DR
KW
     Chloroplast.
FT
     NON TER
                  16
                         16
     SEQUENCE
                16 AA; 1895 MW;
                                 517FE691B89355B9 CRC64;
SQ
  Query Match
                          30.3%; Score 27; DB 8; Length 16;
  Best Local Similarity 15.4%; Pred. No. 1.4e+03;
  Matches
             2; Conservative
                              10; Mismatches 1; Indels
                                                                 0; Gaps
                                                                              0;
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Qу
              :::|:| ::::::
Dh
            3 IRSPEPEVKILVD 15
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RESULT 27 Q8MES8

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ID
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                 PRELIMINARY;
                                   PRT;
                                           16 AA.
AC
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DT
     01-OCT-2002 (TrEMBLrel. 22, Created)
     01-OCT-2002 (TrEMBLrel. 22, Last sequence update)
DΤ
     01-JUN-2003 (TrEMBLrel. 24, Last annotation update)
DT
     PSI P700 apoprotein Al (Fragment).
DE
GN
     PSAA.
os
     Daphniphyllum calycinum.
OG
     Chloroplast.
OC
     Eukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta;
OC
     Spermatophyta; Magnoliophyta; eudicotyledons; core eudicots;
OC
     Saxifragales; Daphniphyllaceae; Daphniphyllum.
OX
     NCBI TaxID=182071;
RN
     [1]
RP
     SEQUENCE FROM N.A.
RA
     Shi S., Huang Y., Zhong Y., Du Y., Zhang Q., Chang H., Boufford D.E.;
RT
     "Phylogeny of the Altingiaceae based on cpDNA matk, PY-IGS and nrDNA
RT
     ITS sequences.";
     Plant Syst. Evol. 230:13-24(2001).
RL
DR
     EMBL; AF377997; AAM45516.1; -.
     GO; GO:0009507; C:chloroplast; IEA.
DR
KW
    Chloroplast.
    NON TER
FT
                  16
                         16
SQ
     SEQUENCE
              16 AA; 1895 MW; 517FE691B89355B9 CRC64;
  Query Match
                          30.3%; Score 27; DB 8; Length 16;
                          15.4%; Pred. No. 1.4e+03;
  Best Local Similarity
             2; Conservative 10; Mismatches
                                                   1; Indels
                                                                 0; Gaps
                                                                              0;
            1 LOTPOPLLOVMME 13
Qу
             :::|:| ::::::
Db
            3 IRSPEPEVKILVD 15
RESULT 28
08MC35
ID
    Q8MC35
                                   PRT:
                                           16 AA.
                 PRELIMINARY;
AC
     Q8MC35;
DT
     01-OCT-2002 (TrEMBLrel. 22, Created)
     01-OCT-2002 (TrEMBLrel. 22, Last sequence update)
DT
     01-JUN-2003 (TrEMBLrel. 24, Last annotation update)
DE
     PsaA (Fragment).
GN
     PSAA.
    Heimia myrtifolia.
OS
OG
    Chloroplast.
OC
     Eukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta;
OC
     Spermatophyta; Magnoliophyta; eudicotyledons; core eudicots; rosids;
    Myrtales; Lythraceae; Heimia.
OC
OX
    NCBI TaxID=135798;
RN
     [1]
RP
     SEQUENCE FROM N.A.
RC
     STRAIN=H491;
RA
     Huang Y., Shi S.;
     "Phylogenetics of Lythraceae sensu lato: A Preliminary Analysis Based
RT
     on Chloroplast rbcL Gene, psaA-ycf3 Spacer, and Nuclear rDNA Internal
RT
RT
     Transcribed Spacer (ITS) Sequences.";
     Int. J. Plant Sci. 163:215-225(2002).
RL
```

```
DR
     EMBL; AY035735; AAL14151.1; -.
DR
     GO; GO:0009507; C:chloroplast; IEA...
KW
     Chloroplast.
FT
     NON TER
                  16
                         16
     SEQUENCE
                16 AA; 1895 MW; 517FE691B89355B9 CRC64;
SQ
  Query Match
                          30.3%; Score 27; DB 8; Length 16;
  Best Local Similarity
                          15.4%; Pred. No. 1.4e+03;
  Matches
             2; Conservative 10; Mismatches
                                                   1: Indels
                                                                  0; Gaps
                                                                              0;
            1 LQTPQPLLQVMME 13
Qy
              :::|:| ::::::
Db
            3 IRSPEPEVKILVD 15
RESULT 29
Q9XQN9
ID
                 PRELIMINARY;
                                   PRT;
                                           17 AA.
     Q9XQN9
AC
     Q9XQN9;
DT
     01-NOV-1999 (TrEMBLrel. 12, Created)
     01-NOV-1999 (TrEMBLrel. 12, Last sequence update)
DT
     01-JUN-2003 (TrEMBLrel. 24, Last annotation update)
DT
DE
     Photosystem I P700 chlorophyll (Fragment).
GN
     PSAA.
OS
     Sinapis alba (White mustard) (Brassica hirta).
OG
     Chloroplast.
OC
     Eukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta;
OC
     Spermatophyta; Magnoliophyta; eudicotyledons; core eudicots; rosids;
OC
     eurosids II; Brassicales; Brassicaceae; Sinapis.
OX
     NCBI TaxID=3728;
RN
     [1]
RP
     SEOUENCE FROM N.A.
RC
     TISSUE=Cotyledon;
    MEDLINE=20136375; PubMed=10672444;
RX
RA
     Summer H., Pfannschmidt T., Link G.;
RT
     "Transcripts and sequence elements suggest differential promoter usage
RT
     within the ycf3-psaAB gene cluster on mustard (Sinapis alba L.)
RT
     chloroplast DNA.";
RL
     Curr. Genet. 37:45-52(2000).
DR
     EMBL; AJ242660; CAB45538.1; -.
     GO; GO:0009507; C:chloroplast; IEA.
DR
KW
     Chloroplast.
     NON TER
                  17
FΤ
                         17
     SEQUENCE
                17 AA; 2008 MW; 99517FE691B89355 CRC64;
SO
  Query Match
                          30.3%;
                                  Score 27; DB 8; Length 17;
  Best Local Similarity
                          15.4%;
                                  Pred. No. 1.5e+03;
  Matches
             2; Conservative
                               10; Mismatches
                                                                              0;
                                                   1; Indels
                                                                  0; Gaps
            1 LQTPQPLLQVMME 13
Qy
              :::|:| ::::::
Db
            3 IRSPEPEVKILVD 15
RESULT 30
090XE2
                 PRELIMINARY;
ID
    Q90XE2
                                   PRT;
                                           17 AA.
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AC
     Q90XE2;
DT
     01-DEC-2001 (TrEMBLrel. 19, Created)
DT
     01-DEC-2001 (TrEMBLrel. 19, Last sequence update)
     01-DEC-2001 (TrEMBLrel. 19, Last annotation update)
DT
DE
     Transforming growth factor beta 4 (Fragment).
     Gallus gallus (Chicken).
OS
OC
     Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC
     Archosauria; Aves; Neognathae; Galliformes; Phasianidae; Phasianinae;
ОÇ
     Gallus.
OX
     NCBI TaxID=9031;
RN
     [1]
     SEQUENCE FROM N.A.
RP
RA
     Pan H.J., Halper J.;
     "5' end sequence of chicken transforming growth factor beta 4.";
RT
     Submitted (JUN-2001) to the EMBL/GenBank/DDBJ databases.
RL
     EMBL; AF395834; AAL05481.1; -.
DR
FT
     NON TER
                  17
                         17
SO
     SEQUENCE
                17 AA; 1721 MW; 5080B3551E71BD63 CRC64;
  Query Match 29.2%; Score 26; DB 13; Length 17; Best Local Similarity 44.4%; Pred. No. 2.1e+03;
  Matches
             4; Conservative
                                 3; Mismatches
                                                   2; Indels
                                                                  0; Gaps
                                                                               0;
            4 POPLLOVMM 12
Qу
              1 111 :::
            3 PSPLLALLL 11
Db
RESULT 31
Q9UCH1
ID
     Q9UCH1
                 PRELIMINARY:
                                    PRT:
                                            16 AA.
AC
     O9UCH1;
DT
     01-MAY-2000 (TrEMBLrel. 13, Created)
     01-MAY-2000 (TrEMBLrel. 13, Last sequence update)
DT
     01-JUN-2003 (TrEMBLrel. 24, Last annotation update)
DT
DΕ
     LIPOAMIDASE (EC 3.1.1.13) (Fragment).
OS
     Homo sapiens (Human).
OC
     Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC
     Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
OX
     NCBI TaxID=9606;
RN
     [1]
RP
     SEQUENCE.
     MEDLINE=93228634; PubMed=8471055;
RX
     Hui D.Y., Hayakawa K., Oizumi J.;
RA
     "Lipoamidase activity in normal and mutagenized pancreatic cholesterol
RT
RT
     esterase (bile salt-stimulated lipase).";
RL
     Biochem. J. 291:65-69(1993).
     GO; GO:0004771; F:sterol esterase activity; IEA.
DR
     SEQUENCE 16 AA; 1844 MW; 147321FA60374B3C CRC64;
SQ
  Query Match
                          28.1%; Score 25; DB 4; Length 16;
                          66.7%; Pred. No. 2.9e+03;
  Best Local Similarity
  Matches
             4; Conservative 1; Mismatches
                                                 1; Indels
                                                                  0; Gaps
                                                                               0;
            1 LOTPOP 6
Qу
              1: 111
Db
            3 LENPQP 8
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RESULT 32
Q9TRB4
ΙD
     Q9TRB4
                 PRELIMINARY;
                                   PRT;
                                           16 AA.
AC
     Q9TRB4;
     01-MAY-2000 (TrEMBLrel. 13, Created)
DT
DT
     01-MAY-2000 (TrEMBLrel. 13, Last sequence update)
DT
     01-JUN-2000 (TrEMBLrel. 14, Last annotation update)
DE
    ATP-dependent 20 S proteasome activator (Fragment).
     Bos taurus (Bovine).
OS
     Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC
OC
    Mammalia; Eutheria; Cetartiodactyla; Ruminantia; Pecora; Bovoidea;
OC
    Bovidae; Bovinae; Bos.
    NCBI TaxID=9913;
OX
RN
     [1]
RP
    SEOUENCE.
RX
    MEDLINE=94342244; PubMed=8063704;
     DeMartino G.N., Moomaw C.R., Zagnitko O.P., Proske R.J., Chu-Ping M.,
RA
     Afendis S.J., Swaffield J.C., Slaughter C.A.;
RA
RT
     "PA700, an ATP-dependent activator of the 20 S proteasome, is an
RT
    ATPase containing multiple members of a nucleotide-binding protein
RT
     family.";
RL
     J. Biol. Chem. 269:20878-20884(1994).
SO
     SEQUENCE
              16 AA; 1878 MW; F70F74211EE26EDE CRC64;
  Query Match
                          28.1%;
                                  Score 25; DB 6; Length 16;
  Best Local Similarity
                          36.4%; Pred. No. 2.9e+03;
 Matches
            4; Conservative
                                 3; Mismatches
                                                  4; Indels
                                                                 0; Gaps
                                                                              0;
Qγ
            4 PQPLLQVMMEP 14
              1:1 1:: 1
Db
            4 PEPNFQLLDNP 14
RESULT 33
O9APT4
    Q9APT4
ID
                 PRELIMINARY;
                                   PRT;
                                           17 AA.
AC
    O9APT4:
DT
     01-JUN-2001 (TrEMBLrel. 17, Created)
DT
     01-JUN-2001 (TrEMBLrel. 17, Last sequence update)
     01-OCT-2003 (TrEMBLrel. 25, Last annotation update)
DT
DE
    Hypothetical protein.
OS
    Pseudomonas aeruginosa.
OC
    Bacteria; Proteobacteria; Gammaproteobacteria; Pseudomonadales;
OC
     Pseudomonadaceae; Pseudomonas.
OX
    NCBI TaxID=287;
RN
     [1]
RP
     SEQUENCE FROM N.A.
RX
    MEDLINE=21142509; PubMed=11208781;
     Liang X., Pham X.Q.T., Olson M.V., Lory S.;
RA
RT
     "Identification of a Genomic Island Present in the Majority of
RT
     Pathogenic Isolates of Pseudomonas aeruginosa.";
RL
     J. Bacteriol. 183:843-853(2001).
DR
    EMBL; AF241171; AAK01539.1; -.
KW
    Hypothetical protein.
SQ
     SEQUENCE
              17 AA; 1954 MW; 68FDF0471E399969 CRC64;
```

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28.1%; Score 25; DB 2; Length 17;
  Query Match
  Best Local Similarity
                          36.4%; Pred. No. 3.1e+03;
             4; Conservative
                                 3: Mismatches
                                                                 0; Gaps
                                                 4; Indels
                                                                              0;
            4 PQPLLQVMMEP 14
Qу
              | |: ::||
Db
            3 PWELVDALLEP 13
RESULT 34
09R5F6
    09R5F6
ID
                 PRELIMINARY;
                                   PRT;
                                           18 AA.
AC
     09R5F6;
     01-MAY-2000 (TrEMBLrel. 13, Created)
DТ
DT
     01-MAY-2000 (TrEMBLrel. 13, Last sequence update)
DΤ
     01-OCT-2003 (TrEMBLrel. 25, Last annotation update)
DE
    Urease small subunit (Fragment).
OS
    Helicobacter mustelae.
    Bacteria; Proteobacteria; Epsilonproteobacteria; Campylobacterales;
OC
OC
    Helicobacteraceae; Helicobacter.
OX
    NCBI TaxID=217;
RN
    [1]
    SEQUENCE.
RP
    MEDLINE=93084378; PubMed=1452359;
RX
    Turbett G.R., Hoj P.B., Horne R., Mee B.J.;
RA
     "Purification and characterization of the urease enzymes of
RT
    Helicobacter species from humans and animals.";
RT
    Infect. Immun. 60:5259-5266(1992).
RL
DR
    PIR; F49215; F49215.
    GO; GO:0016151; F:nickel ion binding; IEA.
DR
DR
     GO; GO:0009039; F:urease activity; IEA.
    GO; GO:0006807; P:nitrogen metabolism; IEA.
DR
     InterPro; IPR002026; Urease gamma.
DR
     Pfam; PF00547; urease gamma; 1.
DR
     SEQUENCE
              18 AA; 2060 MW; 29C8E0AB77E21805 CRC64;
SO
  Query Match
                          28.1%; Score 25; DB 2; Length 18;
  Best Local Similarity
                          33.3%; Pred. No. 3.3e+03;
 Matches
            5; Conservative
                                5; Mismatches
                                                  5; Indels
                                                                 0; Gaps
                                                                             0;
            3 TPQPLLQVMMEPQGD 17
Qу
              ||: | ::|:
                         |:
Db
            2 TPKELDKMMLHYAGE 16
RESULT 35
Q44237
ID
    Q44237
                 PRELIMINARY;
                                   PRT;
                                           11 AA.
AC
    Q44237;
     01-NOV-1996 (TrEMBLrel. 01, Created)
DT
     01-NOV-1996 (TrEMBLrel. 01, Last sequence update)
DT
     01-DEC-2001 (TrEMBLrel. 19, Last annotation update)
DT
DE
    Glutamine synthetase (Fragment).
GN
    GLNA.
OS
    Anabaena sp. (strain PCC 7120).
OC
    Bacteria; Cyanobacteria; Nostocales; Nostocaceae; Nostoc.
```

```
ΟX
    NCBI TaxID=103690;
RN
     [1]
RP
     SEQUENCE FROM N.A.
RC
     STRAIN=PCC 7120;
RA
     Warner L.E., Ligon P.J., Stahel A.W., Curtis S.E.;
RT
     "The apcF gene of Anabaena sp. strain PCC 7120 is regulated by
RT
     nitrogen and the apcF and glnA promoters overlap.";
     Submitted (MAY-1995) to the EMBL/GenBank/DDBJ databases.
RL
RN
    SEQUENCE FROM N.A.
RP
    STRAIN=PCC 7120;
RC
RA
     Scappino L.A.;
RL
     Submitted (FEB-1995) to the EMBL/GenBank/DDBJ databases.
DR
    EMBL; U21853; AAA65652.1; -.
FT
    NON TER
                 11
                         11
     SEQUENCE
SQ
                11 AA; 1316 MW; 2000580E32CB06C7 CRC64;
  Query Match
                          25.8%; Score 23; DB 2; Length 11;
  Best Local Similarity 44.4%; Pred. No. 4.3e+03;
 Matches
             4; Conservative
                                 3; Mismatches
                                                   2; Indels
                                                                 0; Gaps
                                                                             0;
           1 LQTPQPLLQ 9
Qу
             : | | | : | :
Db
           1 MTTPQEVLK 9
RESULT 36
Q9UEE2
ID
    Q9UEE2
                                   PRT;
                 PRELIMINARY;
                                           13 AA.
AC
    Q9UEE2;
     01-MAY-2000 (TrEMBLrel. 13, Created)
DT
DT
     01-MAY-2000 (TrEMBLrel. 13, Last sequence update)
     01-MAY-2000 (TrEMBLrel. 13, Last annotation update)
DT
DE
    ERGB transcription factor (Fragment).
    FLI1.
GN
OS
    Homo sapiens (Human).
OC
    Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC
    Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
OX
    NCBI TaxID=9606;
RN
    [1]
RP
    SEQUENCE FROM N.A.
    Obata K., Hiraga H., Nojima T., Yoshida M.C., Abe S.;
RA
RT
     "Molecular analysis on the breakpoint region of a t(11:22)
RT
    translocation in Ewing's sarcoma.";
    Submitted (MAR-1998) to the EMBL/GenBank/DDBJ databases.
RL
DR
    EMBL; AB012625; BAA32806.1; -.
    NON TER
FT
                  1
                          1
    NON TER
                  13
                         13
FT
    SEQUENCE
                13 AA; 1541 MW; 2C677798CB566AB7 CRC64;
SQ
                          25.8%; Score 23; DB 4; Length 13;
  Query Match
  Best Local Similarity
                         80.0%; Pred. No. 5.1e+03;
 Matches
           4; Conservative
                              0; Mismatches
                                                1; Indels
                                                                 0; Gaps
           2 OTPOP 6
Qу
             1 111
Db
           2 QRPQP 6
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RESULT 37
Q96RT5
ID
                 PRELIMINARY;
                                   PRT;
                                           16 AA.
     Q96RT5
AC
     Q96RT5;
DT
     01-DEC-2001 (TrEMBLrel. 19, Created)
DT
     01-DEC-2001 (TrEMBLrel. 19, Last sequence update)
DT
     01-DEC-2001 (TrEMBLrel. 19, Last annotation update)
DE
     Tuberous sclerosis 1 (Fragment).
GN
     TSC1.
OS
     Homo sapiens (Human).
     Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC
     Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
oc
OX
     NCBI TaxID=9606;
RN
     [1]
RP
     SEQUENCE FROM N.A.
RA
     Fang L., Wu Z.Y., Wang N., MuRong S.X., Lin M.T., Chao D.R.,
RA
     Fang Z.M.;
     Submitted (JUN-2000) to the EMBL/GenBank/DDBJ databases.
RL
DR
     EMBL; AF274227; AAK60414.1; -.
FT
     NON TER
                  1
                         1
     NON TER
                  16
FT
                         16
     SEQUENCE
                16 AA; 1826 MW; 895C250451E3BBEE CRC64;
SO
  Query Match
                          25.8%;
                                  Score 23; DB 4; Length 16;
                          50.0%; Pred. No. 6.3e+03;
  Best Local Similarity
                                                    3; Indels
                                                                  0;
  Matches
             5; Conservative
                                 2; Mismatches
                                                                      Gaps
                                                                              0;
Qу
            1 LQTPQPLLQV 10
              1 11 11::
            7 LNTPGOLLRL 16
Db
RESULT 38
Q84RM4
ID
     Q84RM4
                 PRELIMINARY;
                                   PRT;
                                           16 AA.
AC
     Q84RM4;
ĎΤ
     01-JUN-2003 (TrEMBLrel. 24, Created)
DT
     01-JUN-2003 (TrEMBLrel. 24, Last sequence update)
DT
     01-JUN-2003 (TrEMBLrel. 24, Last annotation update)
DΕ
     Dehydrin-like protein Dh2 (Fragment).
OS
     Boea crassifolia.
OC
     Eukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta;
OC
     Spermatophyta; Magnoliophyta; eudicotyledons; core eudicots; asterids;
     lamiids; Lamiales; Gesneriaceae; Cyrtandroideae; Didymocarpeae; Boea.
OC
OX
     NCBI TaxID=104990;
RN
     [1]
     SEQUENCE FROM N.A.
RP
RA
     Shen Y., Lin Z.;
     "Isolation and structural analysis of a drought-stressed dehydrin-like
RT
RT
     gene BcDh2 promoter from Boea crassifolia.";
     Submitted (FEB-2003) to the EMBL/GenBank/DDBJ databases.
RL
DR
     EMBL; AY243044; AA086689.1; -.
FT
     NON TER
                  16
                         16
SQ
     SEQUENCE
                16 AA; 1887 MW; 399D10A6E3E2ED3E CRC64;
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Query Match
                          25.8%; Score 23; DB 10; Length 16;
  Best Local Similarity
                          66.7%; Pred. No. 6.3e+03;
                                1; Mismatches
 Matches
             4; Conservative
                                                  1; Indels
                                                                 0; Gaps
                                                                              0;
           12 MEPQGD 17
Qу
              111 1:
            1 MEPYGN 6
Db
RESULT 39
Q7X3X0
ID
    Q7X3X0
                 PRELIMINARY;
                                   PRT;
                                           17 AA.
AC
     07X3X0;
DT
     01-OCT-2003 (TrEMBLrel. 25, Created)
     01-OCT-2003 (TrEMBLrel. 25, Last sequence update)
DT
     01-OCT-2003 (TrEMBLrel. 25, Last annotation update)
DE
     Putative histidine kinase (Fragment).
GN
    OSAA.
     Streptomyces lividans.
OS
     Bacteria; Actinobacteria; Actinobacteridae; Actinomycetales;
OC
OC
     Streptomycineae; Streptomycetaceae; Streptomyces.
    NCBI TaxID=1916;
OX
RN
    [1]
     SEOUENCE FROM N.A.
RP
     STRAIN=1326;
RC
     Bishop A., Fielding S., Dyson P., Herron P.;
RA
     "Concerted mutagenesis of a streptomycete genome: identification of a
RT
     novel response regulator involved in osmoadaptation required for
RT
     differentiation and regulation of antibiotic production.";
RT
     Submitted (JUN-2003) to the EMBL/GenBank/DDBJ databases.
RL
RN
    SEQUENCE FROM N.A.
RP
RC
    STRAIN=1326;
RA
     Herron P.R.;
     Submitted (JUN-2003) to the EMBL/GenBank/DDBJ databases.
RL
     EMBL; AJ565873; CAD92482.1; -.
DR
KW
     Kinase.
     NON TER
FΤ
                   1
SQ
     SEQUENCE
                17 AA; 2127 MW; AEA57E24A45AD792 CRC64;
                          25.8%; Score 23; DB 2; Length 17;
  Query Match
                          50.0%; Pred. No. 6.7e+03;
  Best Local Similarity
  Matches
             5; Conservative
                                 1; Mismatches
                                                   4; Indels
                                                                 0; Gaps
                                                                              0;
            4 PQPLLQVMME 13
Qу
              1 11 11 :
Db
            3 PDHLLTVMQQ 12
RESULT 40
Q30216
ID
     Q30216
                 PRELIMINARY;
                                   PRT;
                                           18 AA.
АC
     Q30216;
DT
     01-NOV-1996 (TrEMBLrel. 01, Created)
     01-NOV-1996 (TrEMBLrel. 01, Last sequence update)
DT
     01-JUN-2003 (TrEMBLrel. 24, Last annotation update)
DT
DE
     Human leukocyte antigen alpha chain (Fragment).
```

```
GN
    HLA DQA1.
OS
    Homo sapiens (Human).
OC
     Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
    Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
OC
OX
    NCBI_TaxID=9606;
RN
     [1]
RP
    SEQUENCE FROM N.A.
RX
    MEDLINE=95064785; PubMed=7974465;
RA
    Fogdell A., Olerup O.;
     "The DQA1*0104 allele is carried by DRB1*1001- and DRB1*1401-positive
RT
    haplotypes in Caucasians, Africans and Orientals.";
RT
    Tissue Antigens 44:19-24(1994).
RL
    EMBL; S75685; AAB32621.1; -.
DR
DR
    PIR; I59649; I59649.
FT
    NON TER
                 1
    SEQUENCE
SO
              18 AA; 1708 MW; 097492525C2252FC CRC64;
 Query Match
                         25.8%; Score 23; DB 7; Length 18;
 Best Local Similarity 55.6%; Pred. No. 7.1e+03;
 Matches
            5; Conservative
                                0; Mismatches
                                                 4; Indels
                                                                0; Gaps
                                                                            0;
           8 LQVMMEPQG 16
Qу
             Db
           5 LTTMMSPCG 13
RESULT 41
Q8CG32
ID
    Q8CG32
                                  PRT;
                                          12 AA.
                PRELIMINARY;
AC
    Q8CG32;
DT
    01-MAR-2003 (TrEMBLrel. 23, Created)
    01-MAR-2003 (TrEMBLrel. 23, Last sequence update)
DT
    01-MAR-2003 (TrEMBLrel. 23, Last annotation update)
DE
    L-histidine decarboxylase (Fragment).
    HDC.
GN
OS
    Rattus sp.
OC
    Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC
    Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Rattus.
OX
    NCBI TaxID=10118;
RN
    [1]
RP
    SEQUENCE FROM N.A.
    MEDLINE=96198284; PubMed=8928792;
RX
RA
    Hocker M., Zhang Z., Fenstermacher D.A., Tagerud S., Chulak M.,
    Joseph D., Wang T.C.;
RA
    "Rat histidine decarboxylase promoter is regulated by gastrin through
RT
RT
    a protein kinase C pathway.";
RL
    Am. J. Physiol. 270:G619-G633(1996).
    EMBL; S83368; AAN86715.1; -.
DR
    NON TER
FT
                 12
                        12
    SEQUENCE
               12 AA; 1515 MW; 8783853D28FB5B05 CRC64;
SQ
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 Best Local Similarity 100.0%; Pred. No. 6.9e+03;
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          11 MMEP 14
Qу
              1111
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RESULT 42
088176
ID
    088176
                 PRELIMINARY;
                                   PRT;
                                           13 AA.
     088176;
AC
DT
     01-NOV-1998 (TrEMBLrel. 08, Created)
DT
     01-NOV-1998 (TrEMBLrel. 08, Last sequence update)
     01-OCT-2002 (TrEMBLrel. 22, Last annotation update)
DT
DΕ
    Neural cell adhesion molecule (Fragment).
GN
    NCAM1 OR NCAM.
OS
    Mus musculus (Mouse).
OC
    Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC
    Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Mus.
    NCBI TaxID=10090;
OX
RN
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RP
    SEQUENCE FROM N.A.
RC
     STRAIN=Balb-c; TISSUE=Liver;
    MEDLINE=98250618; PubMed=9582442;
RX
RA
    Kawahigashi H., Harada Y., Asano A., Nakamura M.;
RT
     "A cis-acting regulatory element that affects the alternative splicing
RT
    of a muscle-specific exon in the mouse NCAM gene.";
    Biochim. Biophys. Acta 1397:305-315(1998).
RL
    EMBL; AB001873; BAA31275.1; -.
DR
    MGD; MGI:97281; Ncam1.
DR
    NON TER
FT
                  1
FT
    NON TER
                  13
                         13
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               13 AA; 1496 MW; CC6098E54C72D732 CRC64;
SQ
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                          66.7%; Pred. No. 7.4e+03;
 Matches
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Qу
            4 PQPLLQ 9
              1 1:11
Db
            3 PLPVLQ 8
RESULT 43
O8CIW3
    Q8CIW3
                 PRELIMINARY;
                                   PRT;
ID
                                           13 AA.
AC
    Q8CIW3;
DT
     01-MAR-2003 (TrEMBLrel. 23, Created)
DT
    01-MAR-2003 (TrEMBLrel. 23, Last sequence update)
DT
     01-JUN-2003 (TrEMBLrel. 24, Last annotation update)
DE
    MORF4-related protein (Fragment).
    MORF4L1 OR TEX189.
GN
os
    Mus musculus (Mouse).
OC
    Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC
    Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Mus.
    NCBI_TaxID=10090;
OX
RN
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RP
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RC
    STRAIN=129/SvEv:
    MEDLINE=22220132; PubMed=12234683;
RX
RA
    Tominaga K., Pereira-Smith O.;
```

```
"The genomic organization, promoter position and expression profile of
RT
RT
     the mouse MRG15 gene.";
     Gene 294:215-215(2002).
RL
     EMBL; AY072694; AAL67900.1; -.
DR
    MGD; MGI:1096551; Morf411.
DR
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FT
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                         13
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SO
                13 AA; 1544 MW; 9B0B3B837D5A276A CRC64;
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                                                                 0; Gaps
            2 QTPQPLLQ 9
Qy
              1 1:1
Db
            5 QDPKPKFQ 12
RESULT 44
Q9NNZ2
ID
     Q9NNZ2
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AC
     Q9NNZ2;
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     01-OCT-2000 (TrEMBLrel. 15, Created)
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DT
DT
     01-OCT-2000 (TrEMBLrel. 15, Last annotation update)
     Integrin alpha-2 subunit (Fragment).
DE
GN
     ITGA2.
    Homo sapiens (Human).
OS
OC
    Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
    Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
OC
OX
    NCBI TaxID=9606;
RN
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RP
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    MEDLINE=98421383; PubMed=9746778;
    Kritzik M., Savage B., Nugent D.J., Santoso S., Ruggeri Z.M.,
RA
RA
     Kunicki T.J.;
RT
     "Nucleotide polymorphisms in the alpha2 gene define multiple alleles
RT
     that are associated with differences in platelet alpha2 beta1
RT
    density.";
RL
    Blood 92:2382-2388(1998).
DR
     EMBL; AF062039; AAF77577.1; -.
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FT
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SQ
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  Best Local Similarity
                          71.4%; Pred. No. 9.1e+03;
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            4 PQPLLQV 10
Qу
              I + I + I
           10 PLPLLLV 16
Db
RESULT 45
081CS8
ID
     081CS8
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                 PRELIMINARY:
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DΤ
     01-JUN-2003 (TrEMBLrel. 24, Created)
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01-JUN-2003 (TrEMBLrel. 24, Last sequence update)
DT
     01-JUN-2003 (TrEMBLrel. 24, Last annotation update)
DT
DE
     ThiJ/PfpI family protein.
GN
     BC2668.
     Bacillus cereus (strain ATCC 14579 / DSM 31).
os
     Bacteria; Firmicutes; Bacillales; Bacillaceae; Bacillus.
OC
OX
     NCBI TaxID=226900;
RN
     [1]
RP
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    MEDLINE=22608415; PubMed=12721630;
RX
RA
     Ivanova N., Sorokin A., Anderson I., Galleron N., Candelon B.,
     Kapatral V., Bhattacharyya A., Reznik G., Mikhailova N., Lapidus A.,
RA
     Chu L., Mazur M., Goltsman E., Larsen N., D'Souza M., Walunas T.,
RA
RA
    Grechkin Y., Pusch G., Haselkorn R., Fonstein M., Ehrlich S.D.,
RA
    Overbeek R., Kyrpides N.;
     "Genome sequence of Bacillus cereus and comparative analysis with
RT
RT
     Bacillus anthracis.";
RL
    Nature 423:87-91(2003).
DR
    EMBL; AE017006; AAP09624.1; -.
KW
     Complete proteome.
                18 AA; 2008 MW;
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SO
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                                 2; Mismatches
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                                                                  0; Gaps
            1 LQTPQPL 7
Qу
              1:1 11:
Db
            6 LETVQPV 12
RESULT 46
O9T2Y3
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ID
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AC
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DT
     01-MAY-2000 (TrEMBLrel. 13, Created)
     01-MAY-2000 (TrEMBLrel. 13, Last sequence update)
DT
     01-JUN-2003 (TrEMBLrel. 24, Last annotation update)
DT
DE
     RBCL protein (Fragment).
GN
    RBCL.
OS
    Begonia formosana.
OG
     Chloroplast.
     Eukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta;
OC
OC
     Spermatophyta; Magnoliophyta; eudicotyledons; core eudicots; rosids;
OC
     eurosids I; Cucurbitales; Begoniaceae; Begonia.
     NCBI TaxID=80370;
OX
RN
     [1]
     SEQUENCE FROM N.A.
RP
RA
     Chiang T.Y.;
     "Sequence announcement: rbcL promotor gene in Begonia formosana.";
RT
RL
     Submitted (AUG-1998) to the EMBL/GenBank/DDBJ databases.
     EMBL; AJ009599; CAB52123.1; -.
DR
DR
     EMBL; AJ009598; CAB52121.1; -.
DR
     GO; GO:0009507; C:chloroplast; IEA.
KW
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FT
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SQ
     SEQUENCE
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           12 MEPQGD 17
Qу
              1 MSPQTD 6
Db
RESULT 47
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AC
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     01-OCT-2003 (TrEMBLrel. 25, Created)
DT
DΤ
     01-OCT-2003 (TrEMBLrel. 25, Last sequence update)
DT
     01-OCT-2003 (TrEMBLrel. 25, Last annotation update)
DE
     Phosphorylase kinase (EC 2.7.1.38) (Fragment).
GN
     PHKA1.
OS
    Mus musculus (Mouse).
OC
     Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC
    Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Mus.
OX
    NCBI TaxID=10090;
RN
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RP
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     STRAIN=Balb/C; TISSUE=Skeletal muscle;
RC
    MEDLINE=94043107; PubMed=8226841;
RX
RA
    Wullrich A., Hamacher C., Schneider A., Kilimann M.W.;
RT
     "The multiphosphorylation domain of the phosphorylase kinase alpha M
RT
     alpha L subunits is a hotspot of differential mRNA processing and of
RT
    molecular evolution.";
     J. Biol. Chem. 268:23208-23214(1993).
RL
DR
    EMBL; X73876; CAD92255.1; -.
KW
    Kinase; Transferase.
FT
    NON_TER
                   1
FT
    NON TER
                  10
                         10
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SO
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Db
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RESULT 48
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DT
     01-NOV-1999 (TrEMBLrel. 12, Created)
DT
     01-NOV-1999 (TrEMBLrel. 12, Last sequence update)
DΤ
     01-JUN-2003 (TrEMBLrel. 24, Last annotation update)
DE
    MerT (Fragment).
GN
    MERT.
OS
     Klebsiella oxytoca.
```

```
OC
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OC
     Enterobacteriaceae; Klebsiella.
OX
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RN
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RP
RC
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    MEDLINE=97208220; PubMed=9055422;
RX
    Liebert C.A., Wireman J., Smith T., Summers A.O.;
RA
     "Phylogeny of mercury resistance (mer) operons of gram-negative
RT
    bacteria isolated from the fecal flora of primates.";
RT
    Appl. Environ. Microbiol. 63:1066-1076(1997).
RL
RN
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RP
    SEQUENCE FROM N.A.
RC
    STRAIN=509H;
    MEDLINE=98027386; PubMed=9361435;
RX
RA
    Wireman J., Liebert C.A., Smith T., Summers A.O.;
RT
     "Association of mercury resistance with antibiotic resistance in the
RT
     gram-negative fecal bacteria of primates.";
RL
    Appl. Environ. Microbiol. 63:4494-4503(1997).
RN
     [3]
RP
    SEQUENCE FROM N.A.
RC
    STRAIN=509H;
RA
    Wireman J., Liebert C.A., Smith T., Summers A.O.;
    "15-bp tandem repeats occur at a sharp transition in gc content in the
RT
    mer operon.";
RL
    Submitted (FEB-1999) to the EMBL/GenBank/DDBJ databases.
    EMBL; AF131272; AAD37141.1; -.
DR
DR
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DR
    GO; GO:0015097; F:mercury ion transporter activity; IEA.
DR
    GO; GO:0015694; P:mercury ion transport; IEA.
    InterPro; IPR003457; Transprt MerT.
DR
DR
     Pfam; PF02411; MerT; 1.
FT
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SQ
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Qу
             1 111
            1 MSEPQ 5
Db
RESULT 49
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AC
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     01-NOV-1999 (TrEMBLrel. 12, Created)
DT
     01-NOV-1999 (TrEMBLrel. 12, Last sequence update)
DT
     01-JUN-2003 (TrEMBLrel. 24, Last annotation update)
DT
DE
    MerT (Fragment).
GN
    MERT.
OS
    Escherichia coli.
OC
    Bacteria; Proteobacteria; Gammaproteobacteria; Enterobacteriales;
oc
    Enterobacteriaceae; Escherichia.
OX
    NCBI TaxID=562;
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RN
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RP
RC
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RX
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RA
     "Phylogeny of mercury resistance (mer) operons of gram-negative
RT
RT
     bacteria isolated from the fecal flora of primates.";
RL
     Appl. Environ. Microbiol. 63:1066-1076(1997).
RN
    [2]
RP
     SEQUENCE FROM N.A.
RC
    STRAIN=390;
RX
    MEDLINE=98027386; PubMed=9361435;
    Wireman J., Liebert C.A., Smith T., Summers A.O.;
RA
RT
     "Association of mercury resistance with antibiotic resistance in the
     gram-negative fecal bacteria of primates.";
RT
RL
    Appl. Environ. Microbiol. 63:4494-4503(1997).
RN
RP
    SEQUENCE FROM N.A.
RC
    STRAIN=390;
     Wireman J., Liebert C.A., Smith T., Summers A.O.;
RA
RT
     "15-bp tandem repeats occur at a sharp transition in gc content in the
RT
     mer operon.";
RL
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DR
    EMBL; AF131271; AAD37139.1; -.
    GO; GO:0016020; C:membrane; IEA.
DR
    GO; GO:0015097; F:mercury ion transporter activity; IEA.
DR
    GO; GO:0015694; P:mercury ion transport; IEA.
DR
DR
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FT
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           11 MMEPO 15
              1 111
Db
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DΤ
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\mathsf{DT}
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DT
     01-MAY-2000 (TrEMBLrel. 13, Last annotation update)
DΕ
     Glycoprotein Ib alpha variant D (Fragment).
OS
    Homo sapiens (Human).
OC
     Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
    Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
OC
OX
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RN
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RP
    SEQUENCE FROM N.A.
RX
    MEDLINE=92250564; PubMed=1577776;
    Lopez J.A., Ludwig E.H., McCarthy B.J.;
RA
```

```
RT
    "Polymorphism of human glycoprotein Ib alpha results from a variable
    number of tandem repeats of a 13-amino acid sequence in the mucin-like
RT
RT
    macroglycopeptide region. Structure/function implications.";
RL
    J. Biol. Chem. 267:10055-10061(1992).
DR
    EMBL; S34436; AAB22152.1; -.
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Db
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Search completed: July 4, 2004, 04:45:26 Job time: 32.9254 secs

## GenCore version 5.1.6 Copyright (c) 1993 - 2004 Compugen Ltd.

OM protein - protein search, using sw model

Run on: July 4, 2004, 04:33:26; Search time 5.83582 Seconds

(without alignments)

151.683 Million cell updates/sec

Title: US-09-641-802-2

Perfect score: 89

Sequence: 1 LQTPQPLLQVMMEPQGD 17

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched: 141681 seqs, 52070155 residues

Total number of hits satisfying chosen parameters: 952

Minimum DB seq length: 7
Maximum DB seq length: 18

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 100 summaries

Database: SwissProt 42:\*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

## SUMMARIES

		8					
Result		Query					
No.	Score	Match	Length	DB	ID	Descrip	otion
1	25	28.1	18	1	CPAX BOVIN	P22779	bos taurus
2	23	25.8	15	1	PSAO CUCSA	P42052	cucumis sat
3	22	24.7	14	1	TAT HV1W2	P12509	human immun
4	22	24.7	14	1	TAT HV1Z8	P12511	human immun
5	22	24.7	16	1	CFAB BOVIN	P81187	bos taurus
6	21	23.6	10	1	FAR6 PANRE		panagrellus
7	21	23.6	16	1	FOR1 MYRGU		myrmecia gu
8	21	23.6	18	1	LCTN LAMGL		lama glama
9	20	22.5	11	1	RANC RANPI		rana pipien
10	20	22.5	16	1.	FOR2 MYRGU		myrmecia gu
11	20	22.5	16	1	ODO2 BOVIN		bos taurus
12	19.5	21.9	14	1	UC34 MAIZE	P80640	zea mays (m
13	19	21.3	9	1	AL10 CARMA		carcinus ma
14	19	21.3	9	1	FAR9 ASCSU	P43172	ascaris suu
15	19	21.3	12	1	FIF1 SARBU	P83349	sarcophaga
16	19	21.3	15	1	UC29 MAIZE		zea mays (m
17	19	21.3	17	1	YALA_TRYBB		trypanosoma

18	18	20.2	9	1	OXYT RAJCL	P42994	raja clavat
19	18	20.2	11	1	CEP1 ACHFU		achatina fu
20	18	20.2	13	1	GER1 HORVU		hordeum vul
21	18	20.2	13	1	GER2 HORVU		hordeum vul
22	17	19.1	8	1	WP1 PERAT		perkinsus a
23	17	19.1	9	1	FAR1 CALVO		calliphora
24	17	19.1	9	1	MGMT BOVIN		bos taurus
25	17	19.1	9	1	RT33_BOVIN		bos taurus
26	17	19.1	10	1	RT02 BOVIN		bos taurus
27	17	19.1	11	1	HS70 PINPS		pinus pinas
28	17	19.1	11	1	TKNA GADMO		gadus morhu
29	17	19.1	12	1	TKN1 KASMA		kassina mac
30	17	19.1	13	1	ACT7 SOYBN		glycine max
31	17	19.1	13	ī	LPAA PORGI		porphyromon
32	17	19.1	14	1	ECDC LYMDI		lymantria d
33	17	19.1	15	1	ARCA STRP5		streptococc
34	17	19.1	15	1	URE2 MORMO		morganella
35	17	19.1	15	1	VORA METTM		methanobact
36	17	19.1	17	1	ACT6 SOYBN		glycine max
37	17	19.1	17	1	B29K PORGI		porphyromon
38	17	19.1	17	1	PC24 BRANA		brassica na
39	17	19.1	17	1	PSBL SYNVU		synechococc
40	17	19.1	18	1	HEX ADECU		canine aden
41	16	18.0	8	1	NPMB BOVIN		bos taurus
42	16	18.0	8	1	UF06 MOUSE		mus musculu
43	16	18.0	10	1	TKN PHYBI		phyllomedus
44	16	18.0	10	1	UPA2 HUMAN		homo sapien
45	16	18.0	11	1	BPP3 BOTIN		bothrops in
46	16	18.0	12	1	NUDM CANFA		canis famil
47	16	18.0	12	1	TM2A METMA		methanosarc
48	16	18.0	13	1	CRBL ICASP		icaria sp.
49	16	18.0	14	1	UN37 CLOPA		clostridium
50	16	18.0	15	1	COXI THUOB		thunnus obe
51	16	18.0	15	1	CYSK CLOPA		clostridium
52	16	18.0	15	1	ECDA LYMDI		lymantria d
53	16	18.0	15	1	FKB7 PINPS		pinus pinas
54	16	18.0	15	1	SODM STRGR		streptomyce
55	16	18.0	16	1	BRB BASAL		basella alb
56	16	18.0	16				homo sapien
57	16	18.0	17	1	A45K MYCBO		mycobacteri
58	16	18.0	17	1	APID BOMPA		bombus pasc
59	16	18.0	17	1	EFG THEAQ		thermus aqu
60	16	18.0	18	1	CXA1 CONER		conus ermin
61	16	18.0	18	1	HEMH THETS		theromyzon
62	15	16.9	7	1	CARP MYTED		mytilus edu
63	15	16.9	7	1	UF04 MOUSE		mus musculu
64	15	16.9	8	1	UH09 RAT		rattus norv
65	15	16.9	9	1	SAP STOVA		stomopneute
66	15	16.9	10	1	BPP2 BOTJA		bothrops ja
67	15	16.9	10	1	PNEU HUMAN		homo sapien
68	15	16.9	10	1	PNEU_NOMAN		rattus norv
69	15	16.9	11	1	BPP4 BOTIN		bothrops in
70	15	16.9	11	1	BPPB AGKHA		agkistrodon
71	15	16.9	11	1	TKNA CHICK		gallus gall
72	15	16.9	11	1	TKNA_CHICK		equus cabal
73	15	16.9	11	1	TKNA_NORSE		oncorhynchu
74	15	16.9	11	1	TKNA SCYCA		scyliorhinu
, 1	13	10.3	**	_	IIIIII_SCICA	1333	PCATTOTHILLIN

```
75
         15
              16.9
                        13 1 BP37 LEUMA
                                                          P81754 leucophaea
76
         15
              16.9
                        13 1 BPP1 BOTJA
                                                          P01020 bothrops ja
 77
         15
              16.9
                       13 1
                               EI21 LITRU
                                                         P82097 litoria rub
 78
                      13 1
                               TA13 TREME
         15
              16.9
                                                         P01370 tremella me
 79
                       13 1
                               TY13 PHYRO
         15
              16.9
                                                         P04096 phyllomedus
 80
                        14 1
                               KLPS SCARA
                                                         P58396 scaptocosa
         15
              16.9
 81
         15
              16.9
                        14 1
                               MAST POLJA
                                                         P01517 polistes ja
82
              16.9
                        15 1 IRBP CRISP
         15
                                                         P12665 cricetidae
83
              16.9
         15
                       15 1 LEC3 PSOSC
                                                         P22583 psophocarpu
                       15 1 MAOX CHICK
 84
              16.9
         15
                                                         Q92060 gallus gall
 85
                       15 1 MK1 PALPR
         15
              16.9
                                                         P80408 palomena pr
                       15 1 MK2A_PALPR
 86
         15
              16.9
                                                          P80409 palomena pr
                       15 1 PRP MYCBO
 87
         15
              16.9
                                                         P80149 mycobacteri
                                                 P81106 P83439 metal
P24854 sus scroll
P80410 palomena pr
P80411 palomena pr
P82240 oncorhynchu
P82697 periplaneta
P14444 camelus dro
P14454 lama glama
P15506 bos taurus
P39399 alteromona
Titoria ri
88
         15
              16.9
                       15 1 UC30 MAIZE
                      15 1
89
         15
              16.9
                               UN01 PINPS
90
                               UP02 METAN
         15
              16.9
                      15 1
              16.9
                       16 1
91
         15
                               IBP4 PIG
92
         15
              16.9
                      16 1 MK2B PALPR
                        16 1
93
         15
              16.9
                               MK3 PALPR
                        16 1
                               SAL\overline{3}_ONCMY
94
         15
              16.9
95
         15
              16.9
                       17 1 PH4 PERAM
96
         15
              16.9
                       18 1 FIBA CAMDR
97
         15
              16.9
                       18 1 FIBA LAMGL
98
         15
              16.9
                       18 1 NPMA BOVIN
99
       14.5
              16.3
                       14 1 MARI ALTSP
         14
                        7 1 EIO5 LITRU
100
              15.7
```

## ALIGNMENTS

```
RESULT 1
CPAX BOVIN
    CPAX BOVIN
ID
                    STANDARD;
                                   PRT;
                                           18 AA.
     P22779;
AC
     01-AUG-1991 (Rel. 19, Created)
DT
     01-AUG-1991 (Rel. 19, Last sequence update)
     15-DEC-1998 (Rel. 37, Last annotation update)
DT
DE
    Cytochrome P450 2A (OLF2) (Olfactive) (P52) (EC 1.14.14.1) (Fragment).
OS
     Bos taurus (Bovine).
OC
    Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
    Mammalia; Eutheria; Cetartiodactyla; Ruminantia; Pecora; Bovoidea;
OC
OC
    Bovidae; Bovinae; Bos.
OX
    NCBI TaxID=9913;
RN
    [1]
RP
     SEQUENCE.
RX
    MEDLINE=91027757; PubMed=2121272;
RA
    Lazard D., Tal N., Rubinstein M., Khen M., Lancet D., Zupko K.;
RT
     "Identification and biochemical analysis of novel olfactory-specific
RT
     cytochrome P-450IIA and UDP-glucuronosyl transferase.";
RL
     Biochemistry 29:7433-7440(1990).
     -!- FUNCTION: Cytochromes P450 are a group of heme-thiolate
CC
CC
         monooxygenases. In liver microsomes, this enzyme is involved in an
CC
         NADPH-dependent electron transport pathway. It oxidizes a variety
CC
         of structurally unrelated compounds, including steroids, fatty
CC
         acids, and xenobiotics.
CC
     -!- CATALYTIC ACTIVITY: RH + reduced flavoprotein + O(2) = ROH +
```

```
CC
         oxidized flavoprotein + H(2)0.
CC
     -!- SUBCELLULAR LOCATION: Membrane-bound. Endoplasmic reticulum.
CC
     -!- SIMILARITY: Belongs to the cytochrome P450 family.
     PIR; A35704; A35704.
DR
     InterPro; IPR001128; Cytochrome P450.
DR
     PROSITE; PS00086; CYTOCHROME P450; PARTIAL.
DR
     Oxidoreductase; Monooxygenase; Electron transport; Membrane; Heme;
KW
    Microsome; Endoplasmic reticulum; Olfaction.
KW
     NON TER
                   1
                         1
FT
                                  G \rightarrow D.
FT
    VARIANT
                   6
                          6
    VARIANT
                         11
                                  A -> E.
FT
                  11
     NON TER
FT
                  18
                         18
     SEQUENCE 18 AA; 2058 MW; F80746F76CCD77FF CRC64;
SO
  Query Match
                          28.1%; Score 25; DB 1; Length 18;
  Best Local Similarity
                          46.2%; Pred. No. 4.4e+02;
  Matches
            6; Conservative
                                 0; Mismatches 7; Indels
                                                                 0; Gaps
                                                                             0;
            4 PQPLLQVMMEPQG 16
Qу
             Db
            5 PGPQQQAFKELQG 17
RESULT 2
PSAO CUCSA
     PSAO CUCSA
                                   PRT;
                                           15 AA.
                    STANDARD;
AC
     P42052;
DT
     01-NOV-1995 (Rel. 32, Created)
DT
     01-NOV-1995 (Rel. 32, Last sequence update)
     16-OCT-2001 (Rel. 40, Last annotation update)
DT
     Photosystem 1 reaction centre subunit 8 (Photosystem I 17.5 kDa
DE
DE
     protein) (Fragment).
GN
     PSAM.
OS
     Cucumis sativus (Cucumber).
     Eukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta;
OC
OC
     Spermatophyta; Magnoliophyta; eudicotyledons; core eudicots; rosids;
OC
     eurosids I; Cucurbitales; Cucurbitaceae; Cucumis.
OX
     NCBI TaxID=3659;
RN
    [1]
RP
     SEQUENCE.
RC
     TISSUE=Cotyledon;
     MEDLINE=91355209; PubMed=1883835;
RX
     Iwasaki Y., Ishikawa H., Hibino T., Takabe T.;
RA
     "Characterization of genes that encode subunits of cucumber PS {\tt I}
RT
     complex by N-terminal sequencing.";
RT
     Biochim. Biophys. Acta 1059:141-148(1991).
RL
CC
     -!- FUNCTION: Essential for the activity of NADP photoreduction.
DR
     PIR; E56819; E56819.
KW
     Photosystem I; Photosynthesis; Chloroplast; Thylakoid; Membrane.
FT
     NON TER
                  15
                         15
     SEQUENCE
                15 AA; 1714 MW; CAOBF5DAD403D9F4 CRC64;
                          25.8%; Score 23; DB 1; Length 15;
  Query Match
                          30.8%; Pred. No. 7.8e+02;
  Best Local Similarity
                                3; Mismatches
  Matches
            4; Conservative
                                                6; Indels
                                                                 0; Gaps
                                                                             0:
Qу
```

```
RESULT 3
TAT HV1W2
ID
    TAT HV1W2
                   STANDARD;
                                 PRT; 14 AA.
    P12509;
AC
DΤ
    01-OCT-1989 (Rel. 12, Created)
DT
    01-OCT-1989 (Rel. 12, Last sequence update)
    28-FEB-2003 (Rel. 41, Last annotation update)
DT
DE
    TAT protein (Transactivating regulatory protein) (Fragment).
GN
    TAT.
OS
    Human immunodeficiency virus type 1 (WMJ2 isolate) (HIV-1).
OC
    Viruses; Retroid viruses; Retroviridae; Lentivirus.
OX
    NCBI TaxID=11705;
RN
    [1]
RP
    SEQUENCE FROM N.A.
RX
    MEDLINE=86235450; PubMed=3012778;
    Hahn B.H., Shaw G.M., Taylor M.E., Redfield R.R., Markham P.D.,
RA
    Salahuddin S.Z., Wong-Staal F., Gallo R.C., Parks E.S., Parks W.P.;
RA
RT
    "Genetic variation in HTLV-III/LAV over time in patients with AIDS or
RT
    at risk for AIDS.";
    Science 232:1548-1553(1986).
RL
CC
    -!- FUNCTION: Transcriptional regulator that acts by binding to the
CC
        trans-activating responsive sequence (TAR) RNA element and
CC
        activates transcription initiation and/or elongation from the LTR
CC
        promoter.
CC
    -!- SUBUNIT: Binds cyclin T1 (By similarity).
CC
    -!- SUBCELLULAR LOCATION: Nuclear; nucleolar.
CC
    -!- MISCELLANEOUS: ISOLATES WMJ1, WMJ2, AND WMJ3 WERE OBTAINED FROM
CC
        BLOOD SAMPLES SEQUENTIALLY TAKEN FROM A TWO-YEAR OLD HAITIAN WHO
CC
        WAS PERINATALLY INFECTED BY HER MOTHER.
CC
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    the European Bioinformatics Institute. There are no restrictions on its
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    or send an email to license@isb-sib.ch).
    _____
CC
    EMBL; M12507; AAB12991.1; -.
DR
    HIV; M12507; TAT$WMJ2.
DR
KW
    Transcription regulation; Activator; RNA-binding; Nuclear protein;
    AIDS.
KW
FT
    NON TER
    SEQUENCE 14 AA; 1467 MW; 37CC737BFEF67AA8 CRC64;
SO
  Query Match
                         24.7%; Score 22; DB 1; Length 14;
  Best Local Similarity 60.0%; Pred. No. 1.1e+03;
  Matches
            3; Conservative 2; Mismatches 0; Indels 0; Gaps
                                                                          0;
Qу
          13 EPOGD 17
             :1:11
Db
           4 QPRGD 8
```

```
TAT HV1Z8
    TAT HV1Z8
                 STANDARD; PRT; 14 AA.
    P12511;
AC
DT
    01-OCT-1989 (Rel. 12, Created)
DT
    01-OCT-1989 (Rel. 12, Last sequence update)
DT
    28-FEB-2003 (Rel. 41, Last annotation update)
DE
    TAT protein (Transactivating regulatory protein) (Fragment).
GN
OS
    Human immunodeficiency virus type 1 (Z-84 isolate) (HIV-1).
OC
    Viruses; Retroid viruses; Retroviridae; Lentivirus.
OX
    NCBI TaxID=11681;
RN
    [1]
    SEQUENCE FROM N.A.
RP
    MEDLINE=88281278; PubMed=3395517;
RX
    Yourno J., Josephs S.F., Reitz M.S. Jr., Zagury D., Wong-Staal F.,
RA
RA
    Gallo R.C.;
    "Nucleotide sequence analysis of the env gene of a new Zairian
RT
    isolate of HIV-1.";
RT
    AIDS Res. Hum. Retroviruses 4:165-173(1988).
RL
    -!- FUNCTION: Transcriptional regulator that acts by binding to the
CC
CC
        trans-activating responsive sequence (TAR) RNA element and
CC
        activates transcription initiation and/or elongation from the LTR
CC
        promoter.
CC
    -!- SUBUNIT: Binds cyclin T1 (By similarity).
    -!- SUBCELLULAR LOCATION: Nuclear; nucleolar.
CC
CC
    -!- MISCELLANEOUS: THE Z-84 ISOLATE WAS TAKEN FROM A 54 YEAR-OLD
CC
        ZAIREAN MALE.
CC
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    between the Swiss Institute of Bioinformatics and the EMBL outstation -
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    the European Bioinformatics Institute. There are no restrictions on its
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    or send an email to license@isb-sib.ch).
CC
    _____
DR
    EMBL; J03653; AAA44685.1; -.
DR
    HIV; J03653; TAT$JY1.
KW
    Transcription regulation; Activator; RNA-binding; Nuclear protein;
KW
    AIDS.
    NON TER
FT
    SEQUENCE 14 AA; 1453 MW; 37CC737BF82D7AA8 CRC64;
SO
 Query Match
                        24.7%; Score 22; DB 1; Length 14;
 Best Local Similarity 60.0%; Pred. No. 1.1e+03;
          3; Conservative 2; Mismatches 0; Indels 0; Gaps
                                                                         0;
          13 EPQGD 17
Qv
             :1:11
Db
           4 QPRGD 8
RESULT 5
CFAB BOVIN
ID CFAB BOVIN STANDARD; PRT; 16 AA.
```

RESULT 4

```
AC
     P81187:
     15-JUL-1998 (Rel. 36, Created)
DT
     15-JUL-1998 (Rel. 36, Last sequence update)
DT
     15-MAR-2004 (Rel. 43, Last annotation update)
DT
     Complement factor B (EC 3.4.21.47) (C3/C5 convertase) (EC-VMFB)
DE
DE
     (Fragment).
GN
     BF.
os
     Bos taurus (Bovine).
     Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC
OC
     Mammalia; Eutheria; Cetartiodactyla; Ruminantia; Pecora; Bovoidea;
OC
     Bovidae; Bovinae; Bos.
OX
     NCBI TaxID=9913;
RN
     [1]
     SEOUENCE.
RP
     TISSUE=Blood;
RC
     MEDLINE=97428195; PubMed=9281322;
RX
     Cai G., Satoh T., Hoshi H.;
RA
RT
     "Isolation from fetal bovine serum of a fragment b of complement
RT
     factor B-like protein improving a long-term survival of human
RT
     endothelial cells.";
     Arch. Biochem. Biophys. 345:150-155(1997).
RL
     -!- FUNCTION: Factor B which is part of the alternate pathway of the
CC
CC
         complement system is cleaved by factor D into 2 fragments: Ba and
CC
         Bb. Bb, a serine protease, then combines with complement factor 3b
CC
         to generate the C3 or C5 convertase.
CC
     -!- CATALYTIC ACTIVITY: Cleaves C3 in the alpha-chain to yield C3a and
CC
         C3b. Cleaves C5 in the alpha-chain to yield C5a and C5b. Both
CC
         cleavages take place at the C-terminal of an arginine residue.
CC
     -!- SUBUNIT: Monomer (By similarity).
     -!- SUBCELLULAR LOCATION: Secreted.
CC
     -!- SIMILARITY: Belongs to peptidase family S1.
CC
     InterPro; IPR001254; Peptidase S1.
DR
DR
     PROSITE; PS50240; TRYPSIN DOM; PARTIAL.
     PROSITE; PS00134; TRYPSIN HIS; PARTIAL.
DR
     PROSITE; PS00135; TRYPSIN SER; PARTIAL.
DR
KW
     Complement alternate pathway; Plasma; Hydrolase; Serine protease;
KW
     Glycoprotein; Zymogen.
FT
                        >16
                                  COMPLEMENT FACTOR B BB FRAGMENT.
     CHAIN
                   1
FT
     NON TER
                  16
                         16
SO
     SEOUENCE
              16 AA; 1762 MW; 75FF5D7F5A6A92F0 CRC64;
  Query Match
                          24.7%; Score 22; DB 1; Length 16;
                          25.0%; Pred. No. 1.2e+03;
  Best Local Similarity
  Matches
             2; Conservative
                                 5; Mismatches
                                                   1; Indels
                                                                  0; Gaps
                                                                              0;
            9 QVMMEPQG 16
Qу
              :::::| |
Db
            1 KIVLDPSG 8
RESULT 6
FAR6 PANRE
     FAR6 PANRE
                                   PRT:
                                           10 AA.
ID
                    STANDARD;
AC
     P82660;
DT
     16-OCT-2001 (Rel. 40, Created)
     16-OCT-2001 (Rel. 40, Last sequence update)
DT
     16-OCT-2001 (Rel. 40, Last annotation update)
DT
```

```
DE
     FMRFamide-like neuropeptide PF6 (NGAPQPFVRF-amide).
OS
    Panagrellus redivivus.
OC
    Eukaryota; Metazoa; Nematoda; Chromadorea; Rhabditida;
OC
     Panagrolaimoidea; Panagrolaimidae; Panagrellus.
OX
    NCBI TaxID=6233;
RN
     [1]
    SEQUENCE, FUNCTION, AND AMIDATION.
RP
RA
    Moffett C.L., Marks N.J., Halton D.W., Thomson D.P., Geary T.G.,
RA
    Maule A.G.;
RT
    "Isolation, characterization and pharmacology of RMRFamide-related
    peptides (FaRPs) from free-living nematode, Panagrellus redivivus.";
RT
    Submitted (JUL-2000) to Swiss-Prot.
RL
    -!- FUNCTION: Myoactive.
CC
CC
    -!- SIMILARITY: Belongs to the FARP (FMRFamide related peptide)
CC
         family.
KW
    Neuropeptide; Amidation.
FT
    MOD RES
                  10
                                  AMIDATION.
    SEQUENCE
               10 AA; 1132 MW; CB13E4C9D776C76D CRC64;
SO
 Query Match
                          23.6%; Score 21; DB 1; Length 10;
 Best Local Similarity
                          50.0%; Pred. No. 1.1e+03;
 Matches
             3; Conservative
                                 2; Mismatches
                                                                  0; Gaps
                                                                              0;
                                                 1; Indels
            4 PQPLLQ 9
Qy
              111 ::
            4 POPFVR 9
Db
RESULT 7
FOR1 MYRGU
    FOR1 MYRGU
ΙD
                    STANDARD;
                                   PRT;
                                           16 AA.
AC
    P81438;
    15-DEC-1998 (Rel. 37, Created)
    15-DEC-1998 (Rel. 37, Last sequence update)
DT
    10-OCT-2003 (Rel. 42, Last annotation update)
DT
DE
    Formaecin 1.
OS
    Myrmecia gulosa (Red bulldog ant).
OC
    Eukaryota; Metazoa; Arthropoda; Hexapoda; Insecta; Pterygota;
OC
    Neoptera; Endopterygota; Hymenoptera; Apocrita; Aculeata; Vespoidea;
OC
    Formicidae; Myrmeciinae; Myrmeciini; Myrmecia.
    NCBI TaxID=36170;
OX
RN
RP
    SEQUENCE, AND CARBOHYDRATE-LINKAGE SITE THR-11.
RC
    TISSUE=Hemolymph;
    MEDLINE=98165787; PubMed=9497332;
RX
    Mackintosh J.A., Veal D.A., Beattie A.J., Gooley A.A.;
RA
    "Isolation from an ant Myrmecia gulosa of two inducible
RT
RT
    O-glycosylated proline-rich antibacterial peptides.";
RL
    J. Biol. Chem. 273:6139-6143(1998).
CC
    -!- FUNCTION: Antibacterial peptide. Has activity against E.coli
CC
         but none against other Gram-negative bacteria and Gram-positive
CC
         bacteria.
CC
    -!- SUBCELLULAR LOCATION: Secreted.
CC
    -!- INDUCTION: By bacterial infection.
CC
    -!- PTM: O-LINKED GLYCAN CONSISTS OF A GAL-GALNAC DISACCHARIDE, O-
CC
        GLYCOSYLATION IS ESSENTIAL FOR FULL BIOLOGICAL ACTIVITY.
CC
    -!- SIMILARITY: TO DROSOPHILA DROSOCIN.
```

```
KW
     Antibiotic; Glycoprotein; Insect immunity; Hemolymph.
FT
     CARBOHYD
                  11
                        11
                                  O-LINKED (GALNAC. . .).
SQ
     SEQUENCE 16 AA; 1794 MW; 80CEA3AABBC2E0AE CRC64;
  Query Match
                          23.6%; Score 21; DB 1; Length 16;
                          66.7%; Pred. No. 1.8e+03;
  Best Local Similarity
  Matches
             4; Conservative
                                 0; Mismatches
                                                 2; Indels
                                                                 0; Gaps
                                                                             0;
            3 TPQPLL 8
Qy
              11 TPHPRL 16
Db
RESULT 8
LCTN LAMGL
ID
     LCTN LAMGL
                    STANDARD;
                                   PRT:
                                           18 AA.
AC
     P83315;
DT
     28-FEB-2003 (Rel. 41, Created)
     28-FEB-2003 (Rel. 41, Last sequence update)
DT
     28-FEB-2003 (Rel. 41, Last annotation update)
DT
DE
     Lactophorin (Whey protein) (Fragment).
OS
     Lama glama (Llama).
OC
     Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC
    Mammalia; Eutheria; Cetartiodactyla; Tylopoda; Camelidae; Lama.
OX
    NCBI TaxID=9844;
RN
     [1]
RP
     SEQUENCE.
RX
    MEDLINE=20000588; PubMed=10531593;
     Kappeler S., Farah Z., Puhan Z.;
RA
RT
     "Alternative splicing of lactophorin mRNA from lactating mammary gland
RT
    of the camel (Camelus dromedarius).";
RL
     J. Dairy Sci. 82:1-10(1999).
CC
     -!- SIMILARITY: Belongs to the PP3 / GlyCAM-1 family.
DR
    GO; GO:0005576; C:extracellular; ISS.
DR
    InterPro; IPR007906; GLYCAM-1.
DR
     Pfam; PF05242; GLYCAM-1; 1.
FT
    NON TER
                 18
                        18
     SEQUENCE
                                 CD59616E5B236AC6 CRC64;
SO
               18 AA; 2079 MW;
                          23.6%; Score 21; DB 1; Length 18;
  Query Match
  Best Local Similarity
                         36.4%; Pred. No. 2e+03;
 Matches
            4; Conservative
                                3; Mismatches
                                                  4; Indels
                                                                 0; Gaps
                                                                             0;
Qy
           5 QPLLQVMMEPQ 15
             :| :: || |
Db
           7 EPKDEIYMESQ 17
RESULT 9
RANC RANPI
    RANC RANPI
ID
                   STANDARD;
                                   PRT;
                                           11 AA.
     P08951;
AC
DT
     01-NOV-1988 (Rel. 09, Created)
DT
     01-NOV-1988 (Rel. 09, Last sequence update)
DT
    10-OCT-2003 (Rel. 42, Last annotation update)
DE
    Ranatensin-C.
OS
    Rana pipiens (Northern leopard frog).
```

```
Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC
OC
     Amphibia; Batrachia; Anura; Neobatrachia; Ranoidea; Ranidae; Rana.
OX
     NCBI TaxID=8404;
RN
     [1]
RP
     SEQUENCE.
RC
     TISSUE=Skin secretion;
RX
    MEDLINE=84131098; PubMed=6141890;
ŔĀ
    Nakajima T.;
RL
    Unpublished results, cited by:
RL
     Erspamer V., Erspamer G.F., Mazzanti G., Endean R.;
     Comp. Biochem. Physiol. 77C:99-108(1984).
RL
     -!- SUBCELLULAR LOCATION: Secreted.
CC
     -!- TISSUE SPECIFICITY: Skin.
CC
CC
     -!- SIMILARITY: Belongs to the bombesin/neuromedin B/ranatensin
CC
         family.
DR
     InterPro; IPR000874; Bombesin.
DR
     Pfam; PF02044; Bombesin; 1.
     PROSITE; PS00257; BOMBESIN; 1.
DR
     Amphibian defense peptide; Bombesin family; Amidation.
KW
FT
     MOD RES
                  11
                         11
                                  AMIDATION.
SQ
     SEQUENCE
                11 AA; 1304 MW; D6C9885A61ADC366 CRC64;
  Query Match
                          22.5%; Score 20; DB 1; Length 11;
  Best Local Similarity
                          75.0%; Pred. No. 1.8e+03;
                                                                  0; Gaps
  Matches
             3; Conservative
                                 1; Mismatches
                                                  0; Indels
                                                                              0;
            2 QTPQ 5
Qу
              :111
Db
            1 ZTPO 4
RESULT 10
FOR2 MYRGU
     FOR2 MYRGU
ID
                                   PRT;
                    STANDARD;
                                           16 AA.
AC
     P81437;
DT
     15-DEC-1998 (Rel. 37, Created)
     15-DEC-1998 (Rel. 37, Last sequence update)
DT
    10-OCT-2003 (Rel. 42, Last annotation update)
DE
    Formaecin 2.
os
    Myrmecia gulosa (Red bulldog ant).
oc
     Eukaryota; Metazoa; Arthropoda; Hexapoda; Insecta; Pterygota;
OC
    Neoptera; Endopterygota; Hymenoptera; Apocrita; Aculeata; Vespoidea;
OC
     Formicidae; Myrmeciinae; Myrmeciini; Myrmecia.
OX
     NCBI TaxID=36170;
RN
     [1]
RP
     SEQUENCE, AND CARBOHYDRATE-LINKAGE SITE THR-11.
RC
     TISSUE=Hemolymph;
RX
     MEDLINE=98165787; PubMed=9497332;
RA
    Mackintosh J.A., Veal D.A., Beattie A.J., Gooley A.A.;
RT
     "Isolation from an ant Myrmecia gulosa of two inducible
RT
     O-glycosylated proline-rich antibacterial peptides.";
ŔĹ
     J. Biol. Chem. 273:6139-6143(1998).
CC
     -!- FUNCTION: Antibacterial peptide. Has activity against E.coli but
CC
         none against other Gram-negative bacteria and Gram-positive
CC
         bacteria.
CC
     -!- SUBCELLULAR LOCATION: Secreted.
CC
     -!- INDUCTION: By bacterial infection.
```

```
CC
     -!- PTM: O-LINKED GLYCAN CONSISTS OF A GAL-GALNAC DISACCHARIDE, O-
CC
         GLYCOSYLATION IS ESSENTIAL FOR FULL BIOLOGICAL ACTIVITY.
CC
     -!- SIMILARITY: TO DROSOPHILA DROSOCIN.
KW
     Antibiotic; Glycoprotein; Insect immunity; Hemolymph.
FT
     CARBOHYD
                  11
                                  O-LINKED (GALNAC. . .).
                         11
SO
     SEQUENCE
                16 AA;
                       1807 MW; 9C3CA3B00BC2E0AE CRC64;
  Query Match
                          22.5%; Score 20; DB 1; Length 16;
  Best Local Similarity
                          66.7%; Pred. No. 2.6e+03;
  Matches
             4; Conservative
                                0; Mismatches
                                                  2; Indels
                                                                 0; Gaps
                                                                              0;
            3 TPOPLL 8
Qy
              1111
           11 TPYPRL 16
Db
RESULT 11
ODO2 BOVIN
     ODO2 BOVIN
                    STANDARD;
                                   PRT;
ID
                                           16 AA.
AC
     P11179;
DT
     01-JUL-1989 (Rel. 11, Created)
DT
     01-JUL-1989 (Rel. 11, Last sequence update)
     28-FEB-2003 (Rel. 41, Last annotation update)
DT
     Dihydrolipoamide succinyltransferase component of 2-oxoglutarate
DE
    dehydrogenase complex (EC 2.3.1.61) (E2) (Fragment).
GN
    DLST.
OS
     Bos taurus (Bovine).
     Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC
OC
    Mammalia; Eutheria; Cetartiodactyla; Ruminantia; Pecora; Bovoidea;
     Bovidae; Bovinae; Bos.
OC
OX
    NCBI TaxID=9913;
RN
    [1]
RP
    SEQUENCE.
RC
    TISSUE=Kidney;
RX
    MEDLINE=88005143; PubMed=3115829;
     Bradford A.P., Aitken A., Beg F., Cook K.G., Yeaman S.J.;
RA
RT
     "Amino acid sequence surrounding the lipoic acid cofactor of bovine
RT
     kidney 2-oxoglutarate dehydrogenase complex.";
RL
     FEBS Lett. 222:211-214(1987).
     -!- FUNCTION: The 2-oxoglutarate dehydrogenase complex catalyzes the
CC
CC
         overall conversion of 2-oxoglutarate to succinyl-CoA and CO(2). It
CC
         contains multiple copies of 3 enzymatic components: 2-oxoglutarate
CC
         dehydrogenase (E1), dihydrolipoamide succinyltransferase (E2) and
CC
         lipoamide dehydrogenase (E3).
     -!- CATALYTIC ACTIVITY: Succinyl-CoA + dihydrolipoamide = CoA + S-
CC
CC
         succinyldihydrolipoamide.
CC
     -!- COFACTOR: Contains 1 covalently bound lipoyl cofactor.
CC
     -!- PATHWAY: Tricarboxylic acid cycle.
CC
     -!- SUBUNIT: Forms a 24-polypeptide structural core with octahedral
CC
         symmetry.
CC
     -!- SUBCELLULAR LOCATION: Mitochondrial.
CC
     -!- SIMILARITY: Belongs to the 2-oxoacid dehydrogenase family.
CC
    -!- SIMILARITY: Contains 1 lipoyl-binding domain.
DR
     PIR; S00123; S00123.
DR
    InterPro; IPR003016; Lipoyl BS.
     PROSITE; PS00189; LIPOYL; PARTIAL.
DR
KW
     Tricarboxylic acid cycle; Transferase; Acyltransferase; Lipoyl;
```

```
KW
    Mitochondrion.
FT
    NON TER
             1
                          1
    BINDING
                   5
FT
                          5
                                  LIPOYL.
FΤ
    NON TER
                  16
                         16
SO
    SEQUENCE
               16 AA; 1643 MW; E34CD29B86B3239D CRC64;
 Query Match
                          22.5%;
                                  Score 20; DB 1; Length 16;
  Best Local Similarity
                          50.0%; Pred. No. 2.6e+03;
 Matches
             3; Conservative
                                 1; Mismatches
                                                  2; Indels
                                                                 0; Gaps
                                                                              0;
            1 LQTPQP 6
Qу
              : | | |
            8 VQVPSP 13
Db
RESULT 12
UC34 MAIZE
ID
    UC34 MAIZE
                    STANDARD;
                                   PRT:
                                           14 AA.
АC
    P80640;
    01-OCT-1996 (Rel. 34, Created)
DT
DT
    01-OCT-1996 (Rel. 34, Last sequence update)
DT
    15-MAR-2004 (Rel. 43, Last annotation update)
DΕ
    Unknown protein from 2D-page of etiolated coleoptile (Spot 360)
DΕ
    (Fragments).
os
    Zea mays (Maize).
OC
    Eukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta;
OC
    Spermatophyta; Magnoliophyta; Liliopsida; Poales; Poaceae;
    PACCAD clade; Panicoideae; Andropogoneae; Zea.
OC
    NCBI TaxID=4577;
OX
RN
    [1]
    SEQUENCE.
RP
RC
    TISSUE=Coleoptile;
RA
    Touzet P., Riccardi F., Morin C., Damerval C., Huet J.-C.,
    Pernollet J.-C., Zivy M., de Vienne D.;
RA
RT
     "The maize two dimensional gel protein database: towards an integrated
RT
    genome analysis program.";
    Theor. Appl. Genet. 93:997-1005(1996).
RL
CC
    -!- MISCELLANEOUS: On the 2D-gel the determined pI of this unknown
CC
        protein is: 5.7, its MW is: 41.1 kDa.
CC
    -!- CAUTION: The order of the peptides shown is uncertain.
DR
    Maize-2DPAGE; P80640; COLEOPTILE.
DR
    MaizeDB; 123965; -.
FT
    NON_TER
                   1
FT
    NON_CONS
                   8
                          9
    NON TER
FT
                  14
                         14
    SEQUENCE
SQ
              14 AA; 1527 MW; DC525FF7B0BE682D CRC64;
 Query Match
                          21.9%;
                                  Score 19.5; DB 1; Length 14;
 Best Local Similarity
                         54.5%; Pred. No. 2.8e+03;
 Matches
             6; Conservative
                                 1; Mismatches
                                                   3; Indels
                                                                 1; Gaps
                                                                              1;
            5 QPL-LQVMMEP 14
Qу
              : 11 1 11
```

4 EPLALSVFDEP 14

Db

```
AL10 CARMA
     AL10 CARMA
                                   PRT;
ID
                    STANDARD;
                                            9 AA.
AC
     P81813;
DT
     30-MAY-2000 (Rel. 39, Created)
     30-MAY-2000 (Rel. 39, Last sequence update)
DT
     30-MAY-2000 (Rel. 39, Last annotation update)
DT
DE
     Carcinustatin 10.
     Carcinus maenas (Common shore crab) (Green crab).
OS
     Eukaryota; Metazoa; Arthropoda; Crustacea; Malacostraca;
OC
     Eumalacostraca; Eucarida; Decapoda; Pleocyemata; Brachyura;
OC
     Eubrachyura; Portunoidea; Portunidae; Carcinus.
OC
OX
     NCBI TaxID=6759;
RN
     [1]
     SEQUENCE.
RP
     TISSUE=Cerebral ganglion, and Thoracic ganglion;
RC
     MEDLINE=98121193; PubMed=9461295;
RX
RA
     Duve H., Johnsen A.H., Maestro J.-L., Scott A.G., Jaros P.P.,
RA
     Thorpe A.;
RT
     "Isolation and identification of multiple neuropeptides of the
RT
     allatostatin superfamily in the shore crab Carcinus maenas.";
     Eur. J. Biochem. 250:727-734(1997).
RL
     -!- FUNCTION: May act as a neurotransmitter or neuromodulator.
CC
CC
     -!- SIMILARITY: Belongs to the allatostatin family.
KW
     Neuropeptide; Amidation; Multigene family.
FT
     MOD RES
                                  AMIDATION.
     SEQUENCE
                9 AA; 963 MW; 372D79CDCB4776C7 CRC64;
SQ
  Query Match
                          21.3%; Score 19; DB 1; Length 9;
  Best Local Similarity
                          100.0%; Pred. No. 1.4e+05;
            3; Conservative 0; Mismatches 0; Indels
  Matches
                                                                  0; Gaps
                                                                              0;
            4 POP 6
Qу
              111
            2 PQP 4
Db
RESULT 14
FAR9 ASCSU
     FAR9 ASCSU
                    STANDARD;
                                   PRT:
                                             9 AA.
AC.
     P43172:
DT
     01-NOV-1995 (Rel. 32, Created)
     01-NOV-1995 (Rel. 32, Last sequence update)
DT
     01-FEB-1996 (Rel. 33, Last annotation update)
DT
DE
     FMRFamide-like neuropeptide AF9.
     Ascaris suum (Pig roundworm) (Ascaris lumbricoides).
os
     Eukaryota; Metazoa; Nematoda; Chromadorea; Ascaridida; Ascaridoidea;
OC
OC
     Ascarididae; Ascaris.
OX
     NCBI TaxID=6253;
RN
     [1]
RP
     SEQUENCE.
     MEDLINE=95380362; PubMed=7651904;
RX
     Cowden C., Stretton A.O.W.;
RA
     "Eight novel FMRFamide-like neuropeptides isolated from the nematode
RT
RT
     Ascaris suum.";
RL
     Peptides 16:491-500(1995).
CC
     -!- SIMILARITY: Belongs to the FARP (FMRFamide related peptide)
CC
         family.
```

```
KW
     Neuropeptide; Amidation.
     MOD RES
                   9 9
                                  AMIDATION.
FT
     SEOUENCE
                9 AA; 1012 MW;
                                 524F073774176877 CRC64;
SO
                          21.3%; Score 19; DB 1; Length 9;
  Query Match
                          75.0%; Pred. No. 1.4e+05;
  Best Local Similarity
                                                  0; Indels
 Matches
             3; Conservative
                                 1; Mismatches
                                                                0; Gaps
                                                                               0;
            4 POPL 7
Qу
              1:11
            4 PRPL 7
Db
RESULT 15
FIF1 SARBU
ID
     FIF1 SARBU
                    STANDARD;
                                   PRT;
                                            12 AA.
AC
     P83349;
DT
     28-FEB-2003 (Rel. 41, Created)
     28-FEB-2003 (Rel. 41, Last sequence update) 28-FEB-2003 (Rel. 41, Last annotation update)
DT
DT
    Neb-FIRFamide 1.
DE
     Sarcophaga bullata (Grey flesh fly) (Neobellieria bullata).
os
     Eukaryota; Metazoa; Arthropoda; Hexapoda; Insecta; Pterygota;
OC
     Neoptera; Endopterygota; Diptera; Brachycera; Muscomorpha; Oestroidea;
OC
OC
     Sarcophagidae; Sarcophaga.
     NCBI TaxID=7385;
OX
RN
     [1]
     SEQUENCE, AMIDATION, AND FUNCTION.
RP
RC
     TISSUE=CNS;
RX
     MEDLINE=22342733; PubMed=12438685;
RA
     Meeusen T., Mertens I., Clynen E., Baggerman G., Nichols R.,
     Nachman R.J., Huybrechts R., De Loof A., Schoofs L.;
RA
RT
     "Identification in Drosophila melanogaster of the invertebrate G
     protein-coupled FMRFamide receptor.";
RT
     Proc. Natl. Acad. Sci. U.S.A. 99:15363-15368(2002).
RL
CC
     -!- FUNCTION: Has modulatory actions at skeletal neuromuscular
         junctions.
CC
CC
     -!- SUBCELLULAR LOCATION: Secreted.
CC
     -!- SIMILARITY: Belongs to the FARP (FMRFamide related peptide)
CC
         family.
KW
     Neuropeptide; Amidation.
FT
     MOD RES
                                  AMIDATION.
                  12
                         12
SQ
     SEQUENCE
                12 AA; 1389 MW; 2DC45519C14AB5A7 CRC64;
  Query Match
                          21.3%; Score 19; DB 1; Length 12;
                          100.0%; Pred. No. 2.9e+03;
  Best Local Similarity
             3; Conservative
                               0; Mismatches
                                                  0; Indels
                                                                               0;
            4 PQP 6
Qу
              111
            3 PQP 5
Db
RESULT 16
UC29 MAIZE
                                   PRT;
ID
     UC29 MAIZE
                    STANDARD;
                                            15 AA.
AC
     P80635;
```

```
DT
    01-OCT-1996 (Rel. 34, Created)
DT
    01-OCT-1996 (Rel. 34, Last sequence update)
DT
     15-MAR-2004 (Rel. 43, Last annotation update)
DE
    Unknown protein from 2D-page of etiolated coleoptile (Spot 45)
DE
     (Fragment).
OS
    Zea mays (Maize).
OC
    Eukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta;
OC
    Spermatophyta; Magnoliophyta; Liliopsida; Poales; Poaceae;
OC
    PACCAD clade; Panicoideae; Andropogoneae; Zea.
OX
    NCBI TaxID=4577;
RN
    [1]
    SEQUENCE.
RP
RC
    TISSUE=Coleoptile;
RA
    Touzet P., Riccardi F., Morin C., Damerval C., Huet J.-C.,
RA
    Pernollet J.-C., Zivy M., de Vienne D.;
RT
    "The maize two dimensional gel protein database: towards an integrated
RT
    genome analysis program.";
RL
    Theor. Appl. Genet. 93:997-1005(1996).
CC
    -!- MISCELLANEOUS: On the 2D-gel the determined pI of this unknown
CC
        protein is: 4.9, its MW is: 37.6 kDa.
DR
    Maize-2DPAGE; P80635; COLEOPTILE.
DR
    MaizeDB; 123960; -.
FT
    NON TER
                 1
                         1
    NON TER
                 15
FT
                        15
SQ
    SEQUENCE 15 AA; 1679 MW; 3D53086B16018BC1 CRC64;
 Query Match
                         21.3%; Score 19; DB 1; Length 15;
 Best Local Similarity
                         42.9%; Pred. No. 3.6e+03;
 Matches
           3; Conservative
                               2: Mismatches
                                               2; Indels 0; Gaps
                                                                           0;
Qу
           4 POPLLOV 10
             | ||: :
Db
           6 PIPLVDI 12
RESULT 17
YALA TRYBB
    YALA TRYBB
                   STANDARD;
                                  PRT;
ID
                                          17 AA.
AC
    P17961;
DT
    01-NOV-1990 (Rel. 16, Created)
ĎΤ
    01-NOV-1990 (Rel. 16, Last sequence update)
    16-OCT-2001 (Rel. 40, Last annotation update)
DΤ
DE
    Hypothetical 1.9 kDa protein in aldolase locus (ORFA).
OS
    Trypanosoma brucei brucei.
OC
    Eukaryota; Euglenozoa; Kinetoplastida; Trypanosomatidae; Trypanosoma.
OX
    NCBI TaxID=5702;
RN
    [1]
RP
    SEQUENCE FROM N.A.
RC
    STRAIN=427;
RX
    MEDLINE=90272402; PubMed=2349093;
    Vijayasarathy S., Ernest I., Itzhaki J., Sherman D., Mowatt M.R.,
RA
    Michels P.A.M., Clayton C.E.;
RA
RT
    "The genes encoding fructose bisphosphate aldolase in Trypanosoma
RT
    brucei are interspersed with unrelated genes.";
RL
    Nucleic Acids Res. 18:2967-2975(1990).
    ______
CC
CC
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    the European Bioinformatics Institute. There are no restrictions on its
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    or send an email to license@isb-sib.ch).
CC
    ______
CC
    EMBL; X52586; CAA36821.1; -.
DR
DR
    PIR; S15064; S15064.
KW
    Hypothetical protein.
    SEQUENCE 17 AA; 1939 MW; 5FC2B9AF44AEF420 CRC64;
SO
 Query Match
                        21.3%; Score 19; DB 1; Length 17;
 Best Local Similarity 60.0%; Pred. No. 4.1e+03;
          3; Conservative 1; Mismatches 1; Indels 0; Gaps
                                                                         0;
 Matches
           2 OTPOP 6
Qу
             1 1:1
          11 QLPRP 15
Db
RESULT 18
OXYT RAJCL
    OXYT RAJCL
                  STANDARD;
                                PRT;
                                         9 AA.
AC
    P42994;
DT · 01-NOV-1995 (Rel. 32, Created)
    01-NOV-1995 (Rel. 32, Last sequence update)
DT
    01-NOV-1995 (Rel. 32, Last annotation update)
DE
    Glumitocin.
    Raja clavata (Thornback ray).
OS
    Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Chondrichthyes;
OC
    Elasmobranchii; Squalea; Hypnosqualea; Pristiorajea; Batoidea;
OC
    Rajiformes; Rajidae; Raja.
OX
    NCBI TaxID=7781;
RN
    [1]
RP
    SEOUENCE.
    MEDLINE=66123415; PubMed=5880565;
RX
    Acher R., Chauvet J., Chauvet M.-T., Crepy D.;
RA
RT
    "Phylogeny of neurophyophyseal peptides: isolation of a new hormone,
RT
    glumitocin (Ser 4-Gln 8-ocytocin) present in a cartilaginous fish,
RT
    the ray (Raia clavata).";
    Biochim. Biophys. Acta 107:393-396(1965).
RL
    -!- FUNCTION: ANTIDIURETIC HORMONE.
CC
    -!- SIMILARITY: Belongs to the vasopressin/oxytocin family.
CC
    InterPro; IPR000981; Neurhyp horm.
DR
DR
    Pfam; PF00220; hormone4; 1.
DR
    PROSITE; PS00264; NEUROHYPOPHYS HORM; 1.
KW
    Hormone; Amidation.
FT
    DISULFID
                 1
                        6
                        9
FT
    MOD RES
                 9
                                AMIDATION.
    SEQUENCE 9 AA; 984 MW; 17E9C76EB455B04B CRC64;
                        20.2%; Score 18; DB 1; Length 9;
  Query Match
  Best Local Similarity 100.0%; Pred. No. 1.4e+05;
           3; Conservative 0; Mismatches 0; Indels 0; Gaps
  Matches
                                                                         0:
```

Qу

```
Db 7 PQG 9
```

```
RESULT 19
CEP1 ACHFU
ΙD
     CEP1 ACHFU
                    STANDARD;
                                    PRT:
                                            11 AA.
AC
     P22790;
DT
     01-AUG-1991 (Rel. 19, Created)
DT
     01-AUG-1991 (Rel. 19, Last sequence update)
     01-DEC-1992 (Rel. 24, Last annotation update)
DT
     Cardio-excitatory peptide-1 (ACEP-1).
DE
     Achatina fulica (Giant African snail).
OS
     Eukaryota; Metazoa; Mollusca; Gastropoda; Pulmonata; Stylommatophora;
OC
OC
     Sigmurethra; Achatinoidea; Achatinidae; Achatina.
OX
     NCBI TaxID=6530;
RN
     [1]
RP
     SEQUENCE.
RC
     STRAIN=Ferussac; TISSUE=Heart atrium;
RX
     MEDLINE=90211261; PubMed=2322251;
RA
     Fujimoto K., Ohta N., Yoshida M., Kubota I., Muneoka Y., Kobayashi M.;
RT
     "A novel cardio-excitatory peptide isolated from the atria of the
RT
     African giant snail, Achatina fulica.";
RL
     Biochem. Biophys. Res. Commun. 167:777-783(1990).
CC
     -!- FUNCTION: Potentiates the beat of the ventricle, and has also
CC
         excitatory actions on the penis retractor muscle, the buccal
CC
         muscle and the identified neurons controlling the buccal muscle
CC
         movement of achatina.
CC
     -!- SIMILARITY: TO POSSIBLE PEPTIDE L5 FROM APLYSIA.
DR
     PIR; A34662; A34662.
KW
     Hormone; Amidation.
     MOD RES
FT
                  11
                         11
                                  AMIDATION.
SO
     SEQUENCE
                11 AA; 1305 MW; 82D6D5B9C7741365 CRC64;
  Query Match
                          20.2%; Score 18; DB 1; Length 11;
  Best Local Similarity
                          100.0%; Pred. No. 3.8e+03;
 Matches
             3; Conservative
                                 0; Mismatches
                                                    0; Indels
                                                                   0; Gaps
                                                                               0;
           14 PQG 16
Qy
              111
Db
            7 PQG 9
RESULT 20
GER1 HORVU
     GER1 HORVU
ID
                    STANDARD;
                                    PRT;
                                            13 AA.
     P28525;
AC
DT
     01-DEC-1992 (Rel. 24, Created)
     01-DEC-1992 (Rel. 24, Last sequence update)
DT
     16-OCT-2001 (Rel. 40, Last annotation update)
DT
DΕ
     Germin GS1 (Fragment).
     Hordeum vulgare (Barley).
OS
OC
     Eukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta;
OC
     Spermatophyta; Magnoliophyta; Liliopsida; Poales; Poaceae; Pooideae;
OC
     Triticeae; Hordeum.
OX
     NCBI TaxID=4513;
RN
     [1]
```

```
RP
     SEQUENCE.
     STRAIN=cv. CM 72; TISSUE=Root;
RC
RA
     Hurkman W.J., Tao H.P., Tanaka C.K.;
RT
     "Germin-like polypeptides increase in barley roots during salt
RT
     stress.";
     Plant Physiol. 97:366-374(1991).
RL
     -!- FUNCTION: May play a role in altering the properties of cell
CC
CC
         walls during germinative growth.
CC
     -!- SUBUNIT: Oligomer (believed to be a pentamer but probably hexamer)
CC
         (By similarity).
CC
     -!- SUBCELLULAR LOCATION: SOLUBLE, MICROSOMAL AND CELL WALL FRACTIONS.
     -!- TISSUE SPECIFICITY: Roots and coleoptile. In roots, present in
CC
CC
         the mature region, but not in the tip. Not detected in leaves.
CC
     -!- INDUCTION: Increased by salt stress in roots and decreased by salt
CC
         stress in coleoptile.
CC
     -!- PTM: Glycosylated.
     -!- SIMILARITY: Belongs to the germin family.
CC
DR
     InterPro; IPR001929; Germin.
DR
     PROSITE; PS00725; GERMIN; PARTIAL.
KW
     Apoplast; Cell wall; Glycoprotein; Multigene family.
FT
     UNSURE
                  10
                         10
    NON TER
                  13
                         13
FT
     SEQUENCE
                13 AA; 1470 MW; 43FB588AA3B7B6D7 CRC64;
SQ
                          20.2%; Score 18; DB 1; Length 13;
  Query Match
                          75.0%; Pred. No. 4.5e+03;
  Best Local Similarity
                                0; Mismatches
 Matches
            3; Conservative
                                                  1; Indels
                                                                  0; Gaps
                                                                              0;
            4 PQPL 7
Qу
             1 11
Db
            3 PSPL 6
RESULT 21
GER2 HORVU
     GER2 HORVU
                                   PRT:
ID
                    STANDARD:
                                           13 AA.
     P28526;
AC
     01-DEC-1992 (Rel. 24, Created)
DT
DT
     01-DEC-1992 (Rel. 24, Last sequence update)
DT
     16-OCT-2001 (Rel. 40, Last annotation update)
DE
     Germin GS2 (Fragment).
     Hordeum vulgare (Barley).
OS
     Eukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta;
OC
     Spermatophyta; Magnoliophyta; Liliopsida; Poales; Poaceae; Pooideae;
OC
OC
     Triticeae; Hordeum.
OX
     NCBI TaxID=4513;
RN
     [1]
RP
     SEQUENCE.
     STRAIN=cv. CM 72; TISSUE=Root;
RC
     Hurkman W.J., Tao H.P., Tanaka C.K.;
RA
RT
     "Germin-like polypeptides increase in barley roots during salt
RT
     stress.";
RL
     Plant Physiol. 97:366-374(1991).
CC
     -!- FUNCTION: May play a role in altering the properties of cell walls
CC
         during germinative growth.
CC
     -!- SUBUNIT: Oligomer (believed to be a pentamer but probably hexamer)
CC
         (By similarity).
```

```
CC
     -!- SUBCELLULAR LOCATION: SOLUBLE, MICROSOMAL AND CELL WALL FRACTIONS.
     -!- TISSUE SPECIFICITY: Roots and coleoptile. In roots, present in the
CC
CC
         mature region, but not in the tip. Not detected in leaves.
CC
     -!- INDUCTION: Increased by salt stress in roots and decreased by salt
CC
         stress in coleoptile.
CC
     -!- PTM: Glycosylated.
CC
     -!- SIMILARITY: Belongs to the germin family.
     InterPro; IPR001929; Germin.
DR
     PROSITE; PS00725; GERMIN; PARTIAL.
DR
KW
     Apoplast; Cell wall; Glycoprotein; Multigene family.
FT
     UNSURE
                         10
                  10
     NON TER
FT
                  13
                         13
SQ
     SEQUENCE
               13 AA; 1484 MW; 43FB4A1AA3B7B6D7 CRC64;
                          20.2%; Score 18; DB 1; Length 13;
  Query Match
                         75.0%; Pred. No. 4.5e+03;
  Best Local Similarity
            3; Conservative
                              0; Mismatches
                                                  1; Indels
                                                                 0; Gaps
                                                                             0;
            4 PQPL 7
Qу
              \perp
            3 PSPL 6
Db
RESULT 22
WP1 PERAT
    WP1 PERAT
                                            8 AA.
                    STANDARD;
                                   PRT;
ID
AC
     P83195;
DT
     28-FEB-2003 (Rel. 41, Created)
     28-FEB-2003 (Rel. 41, Last sequence update)
DT
DT
     28-FEB-2003 (Rel. 41, Last annotation update)
DE
     Wall protein-1 (PWP-1) (Fragment).
OS
     Perkinsus atlanticus.
OC
     Eukaryota; Alveolata; Perkinsea; Perkinsida; Perkinsidae; Perkinsus.
OX
     NCBI TaxID=106964;
RN
     [1]
     SEQUENCE, FUNCTION, SUBCELLULAR LOCATION, AND DEVELOPMENTAL STAGE.
RP
RX
     MEDLINE=22044350; PubMed=12049410;
RA
     Montes J.F., Durfort M., Llado A., Garcia-Valero J.;
RT
     "Characterization and immunolocalization of a main proteinaceous
RT
     component of the cell wall of the protozoan parasite Perkinsus
RT
     atlanticus.";
     Parasitology 124:477-484(2002).
RL
     -!- FUNCTION: Is a major protein component of the cell wall. May play
CC
CC
         a key role in the organization of the cell wall and in promoting
CC
         the survival of this parasite.
CC
     -!- SUBCELLULAR LOCATION: Cell wall. Disulfide-linked to other cell
CC
         wall components.
     -!- DEVELOPMENTAL STAGE: Expressed throughout all walled developmental
CC
CC
         stages.
KW
     Cell wall.
FT
     NON TER
                   8
SQ
     SEQUENCE
               8 AA; 765 MW; F1787DD87B1AAB16 CRC64;
                          19.1%; Score 17; DB 1; Length 8;
  Query Match
  Best Local Similarity
                          60.0%; Pred. No. 1.4e+05;
  Matches
             3; Conservative 1; Mismatches 1; Indels
                                                                 0; Gaps
                                                                             0;
```

```
Qу
           12 MEPQG 16
             | \cdot |
Db
            1 MEDEG 5
RESULT 23
FAR1 CALVO
                                   PRT;
     FAR1 CALVO
ID
                    STANDARD;
                                            9 AA.
     P41856;
AC
     01-NOV-1995 (Rel. 32, Created)
DT
DT
     01-NOV-1995 (Rel. 32, Last sequence update)
     01-NOV-1995 (Rel. 32, Last annotation update)
DΤ
DE
     CalliFMRFamide 1.
OS
     Calliphora vomitoria (Blue blowfly).
     Eukaryota; Metazoa; Arthropoda; Hexapoda; Insecta; Pterygota;
OC
OC
     Neoptera; Endopterygota; Diptera; Brachycera; Muscomorpha; Oestroidea;
OC
     Calliphoridae; Calliphora.
OX
     NCBI TaxID=27454;
RN
     [1]
RP
     SEQUENCE.
RC
     TISSUE=Thoracic ganglion;
    MEDLINE=92196111; PubMed=1549595;
RX
RA
     Duve H., Johnsen A.H., Sewell J.C., Scott A.G., Orchard I.,
RA
     Rehfeld J.F., Thorpe A.;
RT
     "Isolation, structure, and activity of -Phe-Met-Arg-Phe-NH2
RT
     neuropeptides (designated calliFMRFamides) from the blowfly
RT
     Calliphora vomitoria.";
RL
     Proc. Natl. Acad. Sci. U.S.A. 89:2326-2330(1992).
CC
     -!- FUNCTION: Able to induce fluid secretion from the isolated
CC
         salivary gland of Calliphora.
CC
    -!- SIMILARITY: Belongs to the FARP (FMRFamide related peptide)
CC
         family.
DR
     PIR; A41978; A41978.
KW
    Neuropeptide; Amidation.
FT
     MOD RES
               9 9
                                  AMIDATION.
SO
     SEQUENCE 9 AA; 1169 MW; 29D00699CAB6C6C7 CRC64;
  Query Match
                          19.1%; Score 17; DB 1; Length 9;
  Best Local Similarity 100.0%; Pred. No. 1.4e+05;
  Matches
            3; Conservative
                                0; Mismatches 0; Indels
                                                              0; Gaps
                                                                             0;
            3 TPQ 5
Qу
              +
            1 TPQ 3
Db
RESULT 24
MGMT BOVIN
     MGMT BOVIN
                    STANDARD;
                                   PRT;
                                            9 AA.
ΙD
     P29177;
AC
     01-DEC-1992 (Rel. 24, Created)
DT
     01-DEC-1992 (Rel. 24, Last sequence update)
DT
DT
     01-OCT-1996 (Rel. 34, Last annotation update)
DE
     Methylated-DNA--protein-cysteine methyltransferase (EC 2.1.1.63) (6-0-
     methylguanine-DNA methyltransferase) (Fragment).
DE
GN
     MGMT.
OS
     Bos taurus (Bovine).
```

```
OC
     Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC
     Mammalia; Eutheria; Cetartiodactyla; Ruminantia; Pecora; Bovoidea;
OC
     Bovidae; Bovinae; Bos.
OX
     NCBI TaxID=9913;
RN
     [1]
RP
     SEQUENCE.
RC
     TISSUE=Thymus;
RX
     MEDLINE=90174912; PubMed=2308822;
     Rydberg B., Hall J., Karran P.;
RT
     "Active site amino acid sequence of the bovine O6-methylquanine-DNA
RT
     methyltransferase.";
    Nucleic Acids Res. 18:17-21(1990).
RL
CC
     -!- FUNCTION: Repair of alkylated guanine in DNA by stoichiometrically
CC
         transferring the alkyl group at the 0-6 position to a cysteine
CC
         residue in the enzyme. This is a suicide reaction: the enzyme is
CC
         irreversibly inactivated.
CC
     -!- CATALYTIC ACTIVITY: DNA (containing 6-0-methylguanine) +
CC
         [protein]-L-cysteine = DNA (without 6-O-methylquanine) + protein
CC
         S-methyl-L-cysteine.
CC
     -!- SIMILARITY: WITH SEGMENTS OF E.COLI ADA AND OGT METHYLTRANSFERASE
CC
         WHICH ENCOMPASS THE ALKYL-ACCEPTOR RESIDUES.
DR
     InterPro; IPR001497; Methyltransf 1.
DR
     PROSITE; PS00374; MGMT; PARTIAL.
     DNA repair; Transferase; Methyltransferase.
KW
    NON TER
FT
                   1
                          1
     ACT SITE
                   9
                          9
FT
                                  ALKYL GROUP ACCEPTOR (BY SIMILARITY).
     NON TER
                   9
FT
                          9
     SEQUENCE 9 AA; 967 MW; 325171A720476047 CRC64;
SQ
                          19.1%; Score 17; DB 1; Length 9;
  Query Match
  Best Local Similarity 60.0%; Pred. No. 1.4e+05;
  Matches
            3; Conservative 1; Mismatches 1; Indels
                                                                 0; Gaps
                                                                              0;
            4 POPLL 8
Qу
              1 1:1
            2 PIPIL 6
Db
RESULT 25
RT33 BOVIN
ID
     RT33 BOVIN
                    STANDARD;
                                   PRT;
                                            9 AA.
AC
     P82926;
DT
     28-FEB-2003 (Rel. 41, Created)
     28-FEB-2003 (Rel. 41, Last sequence update)
     28-FEB-2003 (Rel. 41, Last annotation update)
\mathsf{D}\mathbf{T}
DΕ
    Mitochondrial 28S ribosomal protein S33 (S33mt) (MRP-S33) (Fragment).
GN
    MRPS33.
OS
     Bos taurus (Bovine).
     Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC
     Mammalia; Eutheria; Cetartiodactyla; Ruminantia; Pecora; Bovoidea;
OC
OC
     Bovidae; Bovinae; Bos.
OX
     NCBI TaxID=9913;
RN
    [1]
RP
     SEQUENCE.
RC
     TISSUE=Liver;
RX
     MEDLINE=21276436; PubMed=11279123;
RA
     Koc E.C., Burkhart W., Blackburn K., Moseley A., Spremulli L.L.;
```

```
RT
     "The small subunit of the mammalian mitochondrial ribosome:
RT
    identification of the full complement of ribosomal proteins present.";
RL
    J. Biol. Chem. 276:19363-19374(2001).
CC
    -!- SUBUNIT: Component of the mitochondrial ribosome small subunit
         (28S) which comprises a 12S rRNA and about 30 distinct proteins.
CC
    -!- SUBCELLULAR LOCATION: Mitochondrial.
CC
KW
    Ribosomal protein; Mitochondrion.
FT
    NON TER
                   1
                          1
FT
    NON TER
                   9
                          9
               9 AA; 1032 MW; D341D73776DB05B9 CRC64;
SO
    SEOUENCE
 Query Match
                          19.1%; Score 17; DB 1; Length 9;
  Best Local Similarity
                          57.1%; Pred. No. 1.4e+05;
 Matches
            4; Conservative
                                1; Mismatches
                                                2; Indels
                                                                 0; Gaps
                                                                             0;
           1 LOTPOPL 7
Qу
             1: 111
Dh
           1 LFSEQPL 7
RESULT 26
RT02 BOVIN
    RT02 BOVIN
                                   PRT;
ΙD
                    STANDARD;
                                           10 AA.
AC
     P82923;
DT
    28-FEB-2003 (Rel. 41, Created)
    28-FEB-2003 (Rel. 41, Last sequence update)
DT
    28-FEB-2003 (Rel. 41, Last annotation update)
DT
DE
    Mitochondrial 28S ribosomal protein S2 (MRP-S2) (Fragment).
GN
    MRPS2.
OS
    Bos taurus (Bovine).
OC
    Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
    Mammalia; Eutheria; Cetartiodactyla; Ruminantia; Pecora; Bovoidea;
OC
OC
    Bovidae; Bovinae; Bos.
OX
    NCBI TaxID=9913;
RN
    [1]
RP
    SEOUENCE.
RC
    TISSUE=Liver;
RX
    MEDLINE=21276436; PubMed=11279123;
RA
    Koc E.C., Burkhart W., Blackburn K., Moseley A., Spremulli L.L.;
RT
    "The small subunit of the mammalian mitochondrial ribosome:
RT
    identification of the full complement of ribosomal proteins present.";
    J. Biol. Chem. 276:19363-19374(2001).
RL
CC
    -!- SUBUNIT: Component of the mitochondrial ribosome small subunit
         (28S) which comprises a 12S rRNA and about 30 distinct proteins.
CC
    -!- SUBCELLULAR LOCATION: Mitochondrial.
CC
CC
    -!- SIMILARITY: Belongs to the S2P family of ribosomal proteins.
DR
    InterPro; IPR001865; Ribosomal S2.
     PROSITE; PS00962; RIBOSOMAL S2 1; PARTIAL.
DR
KW
    Ribosomal protein; Mitochondrion.
    NON TER
FT
                   1
                         1
FT
    NON TER
                  10
                         10
SO
    SEQUENCE
               10 AA; 1246 MW; 6A7A6679C04B476B CRC64;
  Query Match
                          19.1%; Score 17; DB 1; Length 10;
                         100.0%; Pred. No. 5.1e+03;
  Best Local Similarity
            3; Conservative 0; Mismatches
                                                                 0; Gaps
 Matches
                                                  0; Indels
                                                                             0;
```

```
12 MEP 14
Qу
              | | | |
Db
            2 MEP 4
RESULT 27
HS70 PINPS
     HS70 PINPS
                                    PRT;
ID
                    STANDARD;
                                            11 AA.
AC
     P81672:
DT
     15-JUL-1999 (Rel. 38, Created)
     15-JUL-1999 (Rel. 38, Last sequence update)
DT
     15-MAR-2004 (Rel. 43, Last annotation update)
DT
DE
     Heat shock 70 kDa protein (Fragment).
OS
     Pinus pinaster (Maritime pine).
OC
     Eukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta;
OC
     Spermatophyta; Coniferopsida; Coniferales; Pinaceae; Pinus.
OX
     NCBI TaxID=71647;
RN
     [1]
RP
     SEQUENCE.
RC
     TISSUE=Needle;
RX
     MEDLINE=99274088; PubMed=10344291;
RA
     Costa P., Pionneau C., Bauw G., Dubos C., Bahrman N., Kremer A.,
RA
     Frigerio J.-M., Plomion C.;
     "Separation and characterization of needle and xylem maritime pine
RT
RT
     proteins.";
RL
     Electrophoresis 20:1098-1108(1999).
     -!- MISCELLANEOUS: On the 2D-gel the determined pI of this protein
CC
CC
         (spot N164) is: 5.4, its MW is: 73 kDa.
CC
     -!- SIMILARITY: Belongs to the heat shock protein 70 family.
KW
     ATP-binding; Heat shock; Multigene family.
FT
     NON TER
                   1
                          1
FT
     NON TER
                  11
                         11
so
     SEOUENCE
                11 AA; 1228 MW; 037C1BE8DAA44DD0 CRC64;
  Query Match
                          19.1%;
                                   Score 17; DB 1; Length 11;
                                  Pred. No. 5.6e+03;
  Best Local Similarity
                          20.0%;
  Matches
             2; Conservative
                                  5; Mismatches
                                                    3; Indels
                                                                   0; Gaps
                                                                               0;
            8 LQVMMEPQGD 17
Qу
              ::::
                     11:
Db
            1 VEIIANDQGN 10
RESULT 28
TKNA GADMO
     TKNA GADMO
ID
                    STANDARD;
                                    PRT;
                                            11 AA.
     P28498;
AC
DT
     01-DEC-1992 (Rel. 24, Created)
     01-DEC-1992 (Rel. 24, Last sequence update)
DT
DT
     10-OCT-2003 (Rel. 42, Last annotation update)
DE
     Substance P.
OS
     Gadus morhua (Atlantic cod).
OC
     Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC
     Actinopteryqii; Neopteryqii; Teleostei; Euteleostei; Neoteleostei;
OC
     Acanthomorpha; Paracanthopterygii; Gadiformes; Gadidae; Gadus.
OX
     NCBI TaxID=8049;
RN
     [1]
```

```
ŘΡ
     SEQUENCE.
RC
     TISSUE=Brain;
RX
     MEDLINE=92298992; PubMed=1376687;
RA
     Jensen J., Conlon J.M.;
RT
     "Substance-P-related and neurokinin-A-related peptides from the brain
     of the cod and trout.";
RT
     Eur. J. Biochem. 206:659-664(1992).
RL
     -!- FUNCTION: Tachykinins are active peptides which excite neurons,
CC
CC
         evoke behavioral responses, are potent vasodilators and
CC
         secretagogues, and contract (directly or indirectly) many smooth
CC
         muscles.
CC
     -!- SUBCELLULAR LOCATION: Secreted.
CC
     -!- SIMILARITY: Belongs to the tachykinin family.
DR
    PIR; S23306; S23306.
DR
    InterPro; IPR002040; Tachy Neurokinin.
     InterPro; IPR008215; Tachykinin.
DR
     Pfam; PF02202; Tachykinin; 1.
DR
DR
    SMART; SM00203; TK; 1.
DR
     PROSITE; PS00267; TACHYKININ; 1.
KW
    Tachykinin; Neuropeptide; Amidation; Neurotransmitter.
FT
    MOD RES
                  11
                         11
                                  AMIDATION (BY SIMILARITY).
     SEQUENCE
                11 AA; 1315 MW; 214860D759D6C6C7 CRC64;
SQ
  Query Match
                                  Score 17; DB 1; Length 11;
                          19.1%;
  Best Local Similarity
                          37.5%; Pred. No. 5.6e+03;
 Matches
             3; Conservative
                                 2; Mismatches
                                                                 0; Gaps
                                                  3; Indels
                                                                              0;
            4 PQPLLQVM 11
Qу
              11 ::1
Db
            4 PQQFIGLM 11
RESULT 29
TKN1 KASMA
     TKN1 KASMA
ID
                    STANDARD;
                                   PRT;
                                           12 AA.
AC
     P08613;
     01-AUG-1988 (Rel. 08, Created)
DT
DT
     01-AUG-1988 (Rel. 08, Last sequence update)
DT
     10-OCT-2003 (Rel. 42, Last annotation update)
DE
    Hylambates kassinin ([Glu2, Pro5] kassinin).
os
     Kassina maculata (African rhacophorid frog) (Hylambates maculatus).
OC
     Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
    Amphibia; Batrachia; Anura; Neobatrachia; Ranoidea; Hyperoliidae;
OC
OC
    Kassina.
OX
    NCBI TaxID=8414;
RN
    [1]
RP
     SEQUENCE.
RC
    TISSUE=Skin secretion;
RA
     Yasuhara T., Nakajima T., Erspamer G.F., Erspamer V.;
     "New tachykinins, Glu2, Pro5-kassinin (hylambates-kassinin) and
RT
RT
     hylambatin, in the skin of the African rhacophorid frog Hylambates
    maculatus.";
RT
RL
     Biomed. Res. 2:613-617(1981).
CC
     -!- FUNCTION: Tachykinins are active peptides which excite neurons,
CC
         evoke behavioral responses, are potent vasodilators and
CC
         secretagogues, and contract (directly or indirectly) many smooth
CC
         muscles.
```

```
-!- SUBCELLULAR LOCATION: Secreted.
CC
    -!- TISSUE SPECIFICITY: Skin.
CC
CC
    -!- SIMILARITY: Belongs to the tachykinin family.
    PIR; S10059; S10059.
DR
    InterPro; IPR002040; Tachy Neurokinin.
DR
DR
    InterPro; IPR008215; Tachykinin.
DR
    Pfam; PF02202; Tachykinin; 1.
DR
    SMART; SM00203; TK; 1.
DR
    PROSITE; PS00267; TACHYKININ; 1.
KW
    Amphibian defense peptide; Tachykinin; Neuropeptide; Amidation.
FT
    MOD RES
                 12
                       12
                               AMIDATION.
    SEQUENCE
               12 AA; 1376 MW; 3E756D279DD6DAB7 CRC64;
SQ
 Query Match
                         19.1%; Score 17; DB 1; Length 12;
  Best Local Similarity
                         60.0%; Pred. No. 6.1e+03;
           3; Conservative 1; Mismatches
                                                               0; Gaps
 Matches
                                               1; Indels
                                                                           0;
Qу
          13 EPQGD 17
             |\cdot|: |
           2 EPKPD 6
Db
RESULT 30
ACT7 SOYBN
    ACT7 SOYBN
                   STANDARD;
                                  PRT;
ID
                                         13 AA.
    P15987;
AC
    01-APR-1990 (Rel. 14, Created)
DT
    01-APR-1990 (Rel. 14, Last sequence update)
DT
    15-JUL-1999 (Rel. 38, Last annotation update)
DT
DE
    Actin 7 (Fragment).
GN
    SAC7.
OS
    Glycine max (Soybean).
    Eukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta;
OC
OC
    Spermatophyta; Magnoliophyta; eudicotyledons; core eudicots; rosids;
    eurosids I; Fabales; Fabaceae; Papilionoideae; Phaseoleae; Glycine.
OC
OX
    NCBI TaxID=3847;
RN
    [1]
RP
    SEQUENCE FROM N.A.
RC
    STRAIN=cv. Wayne;
RX
    MEDLINE=91346640; PubMed=2102831;
RA
    Pearson L., Meagher R.B.;
RT
    "Diverse soybean actin transcripts contain a large intron in the 5'
    untranslated leader: structural similarity to vertebrate muscle actin
RT
RT
    genes.";
RL
    Plant Mol. Biol. 14:513-526(1990).
CC
    -!- FUNCTION: Actins are highly conserved proteins that are involved
CC
        in various types of cell motility and are ubiquitously expressed
CC
        in all eukaryotic cells.
CC
    -!- FUNCTION: Essential component of cell cytoskeleton; plays an
CC
        important role in cytoplasmic streaming, cell shape determination,
CC
        cell division, organelle movement and extension growth.
CC
    -!- SUBCELLULAR LOCATION: Cytoplasmic.
    -!- MISCELLANEOUS: There are at least 16 actin genes in soybean.
CC
    -!- SIMILARITY: Belongs to the actin family.
CC
CC
    ______
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    or send an email to license@isb-sib.ch).
    _____
CC
    EMBL; X17120; CAA34980.1; -.
DR
DR
    PIR; S15755; S15755.
DR
    InterPro; IPR004001; Actin.
    InterPro; IPR004000; Actin like.
DR
    PROSITE; PS00406; ACTINS 1; PARTIAL.
DR
    PROSITE; PS00432; ACTINS_2; PARTIAL.
DR
DR
    PROSITE; PS01132; ACTINS ACT LIKE; PARTIAL.
KW
    Structural protein; Multigene family.
FT
    NON TER
                 13
                       13
    SEQUENCE 13 AA; 1420 MW; 8BEFF3C36D4FD05A CRC64;
SO
 Query Match
                         19.1%; Score 17; DB 1; Length 13;
 Best Local Similarity 75.0%; Pred. No. 6.6e+03;
            3; Conservative
                                              0; Indels 0; Gaps
                               1; Mismatches
                                                                          0;
           5 QPLL 8
Qу
             |||:
           8 QPLV 11
RESULT 31
LPAA PORGI
    LPAA PORGI
ID
                   STANDARD;
                                 PRT;
                                         13 AA.
AC
    P81411;
DT
    15-DEC-1998 (Rel. 37, Created)
    15-DEC-1998 (Rel. 37, Last sequence update)
    15-DEC-1998 (Rel. 37, Last annotation update)
DT
DE
    Lipid-A-associated protein (Fragment).
os
    Porphyromonas gingivalis (Bacteroides gingivalis).
    Bacteria; Bacteroidetes; Bacteroides (class); Bacteroidales;
OC
OC
    Porphyromonadaceae; Porphyromonas.
    NCBI TaxID=837;
OX
RN
    [1]
RP
    SEOUENCE.
RC
    STRAIN=W50;
RX
    MEDLINE=99061194; PubMed=9846737;
RA
    Sharp L., Poole S., Reddi K., Fletcher J., Nair S., Wilson M.,
RA
    Curtis M., Henderson B., Tabona P.;
RT
    "A lipid A-associated protein of Porphyromonas gingivalis, derived
RT
    from the haemagglutinating domain of the RI protease gene family, is
RT
    a potent stimulator of interleukin 6 synthesis.";
RL
    Microbiology 144:3019-3026(1998).
    -!- FUNCTION: IS ASSOCIATED WITH LIPID A, A PHOSPHORYLATED GLYCOLIPID
CC
CC
        THAT ANCHORS THE LIPOPOLYSACCHARIDE TO THE OUTER MEMBRANE OF THE
CC
        CELL.
FT
    VARIANT
                 12
                        12
                                G -> F.
FT
                 13
    NON TER
                       13
SQ
    SEQUENCE 13 AA; 1346 MW; 38EA796EAFA63AB7 CRC64;
  Query Match
                        19.1%; Score 17; DB 1; Length 13;
  Best Local Similarity 100.0%; Pred. No. 6.6e+03;
```

```
0; Indels 0; Gaps
  Matches
             3; Conservative 0; Mismatches
                                                                             0;
           15 QGD 17
Qу
              111
Db
            2 QGD 4
RESULT 32
ECDC LYMDI
     ECDC LYMDI
ID
                    STANDARD;
                                   PRT;
                                           14 AA.
AC
     P80940;
DT
     15-JUL-1998 (Rel. 36, Created)
     15-JUL-1998 (Rel. 36, Last sequence update)
DT
DT
     15-JUL-1998 (Rel. 36, Last annotation update)
     Testis ecdysiotropin peptide C (TE).
DE
OS
     Lymantria dispar (Gypsy moth).
OC
     Eukaryota; Metazoa; Arthropoda; Hexapoda; Insecta; Pterygota;
OC
     Neoptera; Endopterygota; Lepidoptera; Glossata; Ditrysia; Noctuoidea;
OC
     Lymantriidae; Lymantria.
OX
     NCBI_TaxID=13123;
RN
     [1]
RP
     SEQUENCE.
RC
     TISSUE=Brain;
RX
     MEDLINE=97387807; PubMed=9243792;
RA
     Loeb M.J., Wagner R.M., Woods C.W., Gelman D.G., Harrison D.,
RA
     Bell R.A.;
     "Naturally occurring analogs of Lymantria testis ecdysiotropin, a
RT
RT
     gonadotropin isolated from brains of Lymantria dispar pupae.";
     Arch. Insect Biochem. Physiol. 36:37-50(1997).
RL
CC
     -!- FUNCTION: Stimulates synthesis of ecdysteroid in the testes of
CC
         larvae and pupae.
SO
     SEQUENCE
               14 AA; 1553 MW; 17F479531A685CBB CRC64;
                          19.1%; Score 17; DB 1; Length 14;
  Query Match
  Best Local Similarity 75.0%; Pred. No. 7.1e+03;
                                                                 0; Gaps
  Matches
             3: Conservative
                                 0; Mismatches
                                                1; Indels
                                                                             0:
            3 TPQP 6
Qy
             11 1
Db
            8 TPLP 11
RESULT 33
ARCA STRP5
     ARCA STRP5
ID
                    STANDARD;
                                   PRT:
                                           15 AA.
AC
     P58827;
DT
     28-FEB-2003 (Rel. 41, Created)
     28-FEB-2003 (Rel. 41, Last sequence update)
DT
     10-OCT-2003 (Rel. 42, Last annotation update)
DT
     Arginine deiminase (EC 3.5.3.6) (ADI) (Arginine dihydrolase) (AD)
DΕ
DΕ
     (Streptococcal acid glycoprotein) (Fragment).
GN
     ARCA OR SAGP.
os
     Streptococcus pyogenes (serotype M5).
OC
     Bacteria; Firmicutes; Lactobacillales; Streptococcaceae;
OC
     Streptococcus.
OX
     NCBI TaxID=160491;
RN
     [1]
```

```
RP
     SEQUENCE, AND CHARACTERIZATION.
RC
     STRAIN=Manfredo / Serotype M5;
RX
     MEDLINE=98298018; PubMed=9632565;
RA
     Degnan B.A., Palmer J.M., Robson T., Jones C.E.D., Fischer M.,
     Glanville M., Mellor G.D., Diamond A.G., Kehoe M.A., Goodacre J.A.;
RA
RT
     "Inhibition of human peripheral blood mononuclear cell proliferation
     by Streptococcus pyogenes cell extract is associated with arginine
RT
RT
     deiminase activity.";
RL
     Infect. Immun. 66:3050-3058(1998).
CC
     -!- FUNCTION: Antitumor protein. Has a powerful and dose-dependent
CC
         inhibitory effect on antigen, superantigen, or mitogen-stimulated
CC
         human peripheral blood mononuclear cell (PBMC) proliferation. It
CC
         may inhibit cell proliferation by arresting cell cycle and
         inducing apoptosis.
CC
     -!- CATALYTIC ACTIVITY: L-arginine + H(2)0 = L-citrulline + NH(3).
CC
CC
     -!- PATHWAY: Arginine degradation via arginine deiminase; first step.
CC
     -!- SUBUNIT: Homodimer.
CC
     -!- SUBCELLULAR LOCATION: Cytoplasmic (Potential).
CC
     -!- PTM: Glycosylated (By similarity).
CC
     -!- SIMILARITY: Belongs to the arginine deiminase family.
DR
     HAMAP; MF 00242; -; 1.
KW
     Hydrolase; Arginine metabolism; Glycoprotein.
FT
     INIT MET
                  0
                          0
FT
     NON TER
                  15
                         15
     SEQUENCE 15 AA; 1657 MW; D21150201B00EE46 CRC64;
SO
  Query Match
                          19.1%; Score 17; DB 1; Length 15;
  Best Local Similarity
                          100.0%; Pred. No. 7.7e+03;
  Matches
            3; Conservative
                              0; Mismatches
                                                  0; Indels
                                                                 0; Gaps
                                                                              0;
            2 QTP 4
Qу
              \perp
Db
            3 OTP 5
RESULT 34
URE2 MORMO
ID
     URE2 MORMO
                                   PRT;
                    STANDARD;
                                           15 AA.
AC
     P17338;
     01-AUG-1990 (Rel. 15, Created)
DT
DT
     01-AUG-1990 (Rel. 15, Last sequence update)
DT
     15-MAR-2004 (Rel. 43, Last annotation update)
     Urease beta subunit (EC 3.5.1.5) (15 kDa subunit) (Urea
DE
DE
     amidohydrolase) (Fragment).
    UREB.
GN
OS
    Morganella morganii (Proteus morganii).
OC
     Bacteria; Proteobacteria; Gammaproteobacteria; Enterobacteriales;
OC
     Enterobacteriaceae; Morganella.
OX
    NCBI TaxID=582;
RN
     [1]
RP
     SEQUENCE.
RX
    MEDLINE=90264298; PubMed=2345135;
RA
     Hu L.-T., Nicholson E.B., Jones B.D., Lynch M.J., Mobley H.L.T.;
RT
     "Morganella morganii urease: purification, characterization, and
RT
     isolation of gene sequences.";
RL
     J. Bacteriol. 172:3073-3080(1990).
CC
     -! - CATALYTIC ACTIVITY: Urea + H(2)0 = CO(2) + 2 NH(3).
```

```
CC
     -!- SUBUNIT: (Alpha, beta, gamma)(3) (By similarity).
     -!- SIMILARITY: Belongs to the urease beta subunit family.
CC
     PIR; B35389; B35389.
DR
KW
     Hydrolase.
     NON TER
FT
                  15
                         15
     SEQUENCE
               15 AA; 1530 MW; 2D98944F2F20C7E8 CRC64;
SO
  Query Match
                          19.1%; Score 17; DB 1; Length 15;
                          75.0%; Pred. No. 7.7e+03:
  Best Local Similarity
                                0; Mismatches
  Matches
            3; Conservative
                                                  1; Indels
                                                              0; Gaps
                                                                             0;
Qу
            4 POPL 7
             1 11
            6 PTPL 9
Db
RESULT 35
VORA METTM
ID
     VORA METTM
                    STANDARD;
                                   PRT;
                                           15 AA.
     P80907;
AC
     01-NOV-1997 (Rel. 35, Created)
DT
DT
     01-NOV-1997 (Rel. 35, Last sequence update)
DT
     15-MAR-2004 (Rel. 43, Last annotation update)
     Ketoisovalerate oxidoreductase subunit vorA (EC 1.-.-.) (VOR) (2-
DE
     oxoisovalerate oxidoreductase alpha chain) (2-oxoisovalerate-
DΕ
DΕ
     ferredoxin oxidoreductase alpha subunit) (Fragment).
GN
OS
    Methanobacterium thermoautotrophicum (strain Marburg / DSM 2133).
OC
    Archaea; Euryarchaeota; Methanobacteria; Methanobacteriales;
OC
    Methanobacteriaceae; Methanothermobacter.
OX
     NCBI TaxID=79929;
RN
     [1]
RP
     SEQUENCE.
RX
     MEDLINE=97261844; PubMed=9108258;
     Tersteegen A., Linder D., Thauer R.K., Hedderich R.;
RT
     "Structures and functions of four anabolic 2-oxoacid oxidoreductases
RT
     in Methanobacterium thermoautotrophicum.";
RL
     Eur. J. Biochem. 244:862-868(1997).
CC
     -!- SUBUNIT: HETEROTRIMER OF THE VORA, VORB AND VORC SUBUNITS.
CC
     -!- MISCELLANEOUS: Has a pH optimum of 9.7 and an optimal temperature
CC
         of 75 degrees Celsius.
KW
     Oxidoreductase.
FT
     NON TER
SQ
     SEQUENCE
               15 AA; 1779 MW; 31320B6531CA528F CRC64;
  Query Match
                          19.1%;
                                 Score 17; DB 1; Length 15;
  Best Local Similarity
                         30.0%; Pred. No. 7.7e+03;
  Matches
            3; Conservative
                              3; Mismatches
                                                 4; Indels
                                                                 0; Gaps
                                                                             0;
Qу
            1 LOTPOPLLOV 10
             :: | | :|
            5 IRKPDSLXEV 14
Db
RESULT 36
ACT6 SOYBN
ID ACT6 SOYBN
                   STANDARD; PRT;
                                           17 AA.
```

```
AC
    P15986;
DT
    01-APR-1990 (Rel. 14, Created)
    01-APR-1990 (Rel. 14, Last sequence update)
    15-JUL-1999 (Rel. 38, Last annotation update)
DT
DE
    Actin 6 (Fragment).
GN
    SAC6.
os
    Glycine max (Soybean).
OC
    Eukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta;
OC
    Spermatophyta; Magnoliophyta; eudicotyledons; core eudicots; rosids;
    eurosids I; Fabales; Fabaceae; Papilionoideae; Phaseoleae; Glycine.
OC
OX
    NCBI TaxID=3847;
RN
    [1]
    SEQUENCE FROM N.A.
RP
RC
    STRAIN=cv. Wayne;
RX
    MEDLINE=91346640; PubMed=2102831;
    Pearson L., Meagher R.B.;
    "Diverse soybean actin transcripts contain a large intron in the 5'
RT
RT
    untranslated leader: structural similarity to vertebrate muscle actin
RT
RL
    Plant Mol. Biol. 14:513-526(1990).
CC
    -!- FUNCTION: Actins are highly conserved proteins that are involved
        in various types of cell motility and are ubiquitously expressed
CC
CC
        in all eukaryotic cells.
CC
    -!- FUNCTION: Essential component of cell cytoskeleton; plays an
CC
        important role in cytoplasmic streaming, cell shape determination,
CC
        cell division, organelle movement and extension growth.
CC
    -!- SUBCELLULAR LOCATION: Cytoplasmic.
CC
    -!- MISCELLANEOUS: There are at least 16 actin genes in soybean.
CC
    -!- SIMILARITY: Belongs to the actin family.
CC
CC
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    between the Swiss Institute of Bioinformatics and the EMBL outstation -
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    the European Bioinformatics Institute. There are no restrictions on its
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    or send an email to license@isb-sib.ch).
    _____
CC
DR
    EMBL; X17119; CAA34979.1; -.
DR
    PIR; S15754; S15754.
DR
    InterPro; IPR004001; Actin.
    InterPro; IPR004000; Actin_like.
DR
    PROSITE; PS00406; ACTINS_1; PARTIAL.
DR
    PROSITE; PS00432; ACTINS_2; PARTIAL.
DR
    PROSITE; PS01132; ACTINS ACT LIKE; PARTIAL.
DR
    Structural protein; Multigene family.
KW
\mathbf{FT}
    NON TER
                 17
                       17
    SEQUENCE 17 AA; 1749 MW; 38F4970D4BEFF3C3 CRC64;
SO
                         19.1%; Score 17; DB 1; Length 17;
  Query Match
  Best Local Similarity 75.0%; Pred. No. 8.7e+03;
  Matches
          3; Conservative 1; Mismatches 0; Indels 0; Gaps
                                                                          0;
Qy
           5 OPLL 8
             111:
           8 QPLV 11
Db
```

```
RESULT 37
B29K PORGI
ID
     B29K PORGI
                    STANDARD;
                                   PRT:
                                           17 AA.
АC
     P81784;
     15-JUL-1999 (Rel. 38, Created)
DT
     15-JUL-1999 (Rel. 38, Last sequence update)
DT
     30-MAY-2000 (Rel. 39, Last annotation update)
DT
     29 kDa immunogenic protein (Fragment).
DE
     Porphyromonas gingivalis (Bacteroides gingivalis).
OS
OC
     Bacteria; Bacteroidetes; Bacteroides (class); Bacteroidales;
OC
     Porphyromonadaceae; Porphyromonas.
OX
     NCBI TaxID=837;
RN
     [1]
RP
     SEQUENCE.
RC
     STRAIN=VPB 3547;
RX
    MEDLINE=20198497; PubMed=10731616;
RA
    Norris J.M., Love D.N.;
RT
     "Serum antibody responses of cats to soluble whole cell antigens of
RT
     feline Porphyromonas gingivalis.";
     Vet. Microbiol. 73:37-49(2000).
RL
KW
    Antigen.
FT
     NON TER
                  17
                         17
     SEQUENCE
                17 AA; 1756 MW; 383DE6AA14331AE0 CRC64;
SQ
  Query Match
                          19.1%; Score 17; DB 1; Length 17;
                          100.0%; Pred. No. 8.7e+03;
  Best Local Similarity
             3; Conservative 0; Mismatches 0; Indels
                                                                  0; Gaps
                                                                              0;
           15 QGD 17
Qу
              +111
           14 QGD 16
Db
RESULT 38
PC24 BRANA
     PC24 BRANA
                                           17 AA.
ID
                    STANDARD;
                                   PRT:
     P81097;
AC
DT
     15-DEC-1998 (Rel. 37, Created)
DT
     15-DEC-1998 (Rel. 37, Last sequence update)
DT
     15-DEC-1998 (Rel. 37, Last annotation update)
DE
     20 kDa pollen coat protein (Fragment).
os
     Brassica napus (Rape).
OC
     Eukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta;
OC
     Spermatophyta; Magnoliophyta; eudicotyledons; core eudicots; rosids;
     eurosids II; Brassicales; Brassicaceae; Brassica.
OC
OX
     NCBI TaxID=3708;
RN
     [1]
RP
     SEQUENCE.
RC
     STRAIN=cv. Topas; TISSUE=Pollen;
     MEDLINE=98345939; PubMed=9680961;
RX
RA
     Murphy D.J., Ross J.H.E.;
RT
     "Biosynthesis, targeting and processing of oleosin-like proteins,
RT
     which are major pollen coat components in Brassica napus.";
RL
     Plant J. 13:1-16(1998).
CC
     -!- FUNCTION: Major component of the pollen coat.
CC
     -!- TISSUE SPECIFICITY: Pollen.
```

```
CC
     -!- SIMILARITY: Belongs to the oleosin family.
     InterPro; IPR000136; Oleosin.
DR
DR
     PROSITE; PS00811; OLEOSINS; PARTIAL.
FT
     NON TER
                 17
                         17
     SEQUENCE
               17 AA; 2018 MW; DC593833F02C52C9 CRC64;
SQ
  Query Match
                          19.1%; Score 17; DB 1; Length 17;
                         75.0%; Pred. No. 8.7e+03;
  Best Local Similarity
  Matches
            3; Conservative
                                1; Mismatches
                                                0; Indels
                                                                 0; Gaps
                                                                             0;
            1 LQTP 4
Qу
              11:1
            1 LQSP 4
Db
RESULT 39
PSBL SYNVU
ID
     PSBL SYNVU
                    STANDARD;
                                   PRT:
                                          17 AA.
     P12241;
AC
     01-OCT-1989 (Rel. 12, Created)
DT
     01-OCT-1989 (Rel. 12, Last sequence update)
DT
     10-OCT-2003 (Rel. 42, Last annotation update)
DT
     Photosystem II reaction center L protein (PSII 5 kDa protein)
DE
DE
    (Fragment).
GN
    PSBL.
    Synechococcus vulcanus (Thermosynechococcus vulcanus).
OS
OC
     Bacteria; Cyanobacteria; Chroococcales; Thermosynechococcus.
OX
    NCBI TaxID=32053;
RN
    [1]
    SEQUENCE.
RP
     Ikeuchi M., Koike H., Inoue Y.;
RA
RT
     "Identification of psbI and psbL gene products in cyanobacterial
RT
     photosystem II reaction center preparation.";
    FEBS Lett. 251:155-160(1989).
RL
    -!- FUNCTION: Not known, it is however required for PSII activity.
CC
CC
     -!- SUBCELLULAR LOCATION: Cellular thylakoid membrane.
    -!- SIMILARITY: Belongs to the psbL family.
CC
    PIR; S05033; S05033.
DR
DR
    HAMAP; MF 01317; -; 1.
DR
    InterPro; IPR003372; PSII PsbL.
DR
    Pfam; PF02419; PsbL; 1.
KW
     Photosynthesis; Thylakoid; Photosystem II; Reaction center.
     NON TER
FT
                 17
                        17
     SEOUENCE
               17 AA; 2018 MW; A86FD372B70B9847 CRC64;
SO
  Query Match
                          19.1%; Score 17; DB 1; Length 17;
  Best Local Similarity
                         100.0%; Pred. No. 8.7e+03;
            3; Conservative
                                                                 0; Gaps
                                0; Mismatches
                                                   0; Indels
                                                                             0;
           12 MEP 14
Qy
              H
           1 MEP 3
Db
RESULT 40
HEX ADECU
ID HEX ADECU
                   STANDARD;
                                 PRT;
                                          18 AA.
```

```
AC
    P35985;
DT
    01-JUN-1994 (Rel. 29, Created)
DT
    01-JUN-1994 (Rel. 29, Last sequence update)
    01-NOV-1997 (Rel. 35, Last annotation update)
DT
DE
    Hexon protein (Late protein 2) (Fragment).
GN
os
    Canine adenovirus type 1 (strain Utrecht).
OC
    Viruses; dsDNA viruses, no RNA stage; Adenoviridae; Mastadenovirus.
    NCBI TaxID=36364;
OX
RN
    [1]
RP
    SEQUENCE FROM N.A.
RX
    MEDLINE=93033182; PubMed=1413543;
    Cai F., Weber J.M.;
RA
RT
    "Nucleotide and deduced amino acid sequence of the canine adenovirus
RT
    type 1 proteinase.";
RL
    Virus Genes 6:307-312(1992).
CC
    -!- FUNCTION: This protein is one of the structural proteins in the
CC
        viral coat and is synthesized during late infection.
    -!- SUBUNIT: Homotrimer (By similarity).
CC
    ______
CC
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    between the Swiss Institute of Bioinformatics and the EMBL outstation -
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    the European Bioinformatics Institute. There are no restrictions on its
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    ______
DR
    EMBL; M72715; AAA42528.1; -.
DR
    PIR; A48550; A48550.
DR
    HSSP; P03277; 1DHX.
KW
    Coat protein; Hexon protein; Late protein.
FT
    NON TER
              1 1
    SEQUENCE 18 AA; 1938 MW; B9E2AE307A420962 CRC64;
SO
                       19.1%; Score 17; DB 1; Length 18;
 Query Match
 Best Local Similarity 75.0%; Pred. No. 9.2e+03;
 Matches 3; Conservative 1; Mismatches 0; Indels 0; Gaps
Qу
          1 LQTP 4
            1:11
Db
          7 LRTP 10
RESULT 41
NPMB BOVIN
    NPMB BOVIN
                  STANDARD;
                                PRT;
    P15507;
AC
    01-APR-1990 (Rel. 14, Created)
DT
    01-APR-1990 (Rel. 14, Last sequence update)
DΤ
    10-OCT-2003 (Rel. 42, Last annotation update)
DT
DE
    Morphine modulating neuropeptide B.
os
    Bos taurus (Bovine).
OC
    Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC
    Mammalia; Eutheria; Cetartiodactyla; Ruminantia; Pecora; Bovoidea;
OC
    Bovidae; Bovinae; Bos.
OX
    NCBI TaxID=9913;
```

```
RN
     [1]
RP
     SEQUENCE, AND SYNTHESIS.
RC
     TISSUE=Brain;
RX
    MEDLINE=86067985; PubMed=3865193;
     Yang H.-Y.T., Fratta W., Majane E.A., Costa E.;
RA
RT
     "Isolation, sequencing, synthesis, and pharmacological
RT
     characterization of two brain neuropeptides that modulate the action
RT
     of morphine.";
RL
     Proc. Natl. Acad. Sci. U.S.A. 82:7757-7761(1985).
CC
     -!- FUNCTION: Modulates the action of morphine.
CC
     -!- SUBCELLULAR LOCATION: Secreted.
     PIR; B24749; B24749.
DR
KW
     Neuropeptide; Amidation.
    MOD RES
FT
                   8
                          8
                                  AMIDATION.
                8 AA; 1082 MW;
     SEQUENCE
SQ
                                 87D416C776D9C729 CRC64;
  Query Match
                          18.0%; Score 16; DB 1; Length 8;
  Best Local Similarity
                          40.0%; Pred. No. 1.4e+05;
                                 2; Mismatches
  Matches
             2; Conservative
                                                 1; Indels
                                                                 0;
                                                                     Gaps
                                                                              0;
           11 MMEPQ 15
Qу
             : : | |
            2 LFQPQ 6
Db
RESULT 42
UF06 MOUSE
    UF06 MOUSE
ID
                    STANDARD;
                                   PRT;
                                            8 AA.
     P38644;
AC
DT
     01-OCT-1994 (Rel. 30, Created)
DT
     01-OCT-1994 (Rel. 30, Last sequence update)
     15-MAR-2004 (Rel. 43, Last annotation update)
DT
DE
     Unknown protein from 2D-page of fibroblasts (P50) (Fragment).
OS
    Mus musculus (Mouse).
OC
     Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
    Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Mus.
OC
OX
    NCBI TaxID=10090;
RN
    [1]
RP
     SEQUENCE.
RC
     TISSUE=Fibroblast;
RX
    MEDLINE=95009907; PubMed=7523108;
    Merrick B.A., Patterson R.M., Wichter L.L., He C., Selkirk J.K.;
RA
     "Separation and sequencing of familiar and novel murine proteins
RT
RT
     using preparative two-dimensional gel electrophoresis.";
     Electrophoresis 15:735-745(1994).
RL
     -!- MISCELLANEOUS: On the 2D-gel the determined pI of this unknown
CC
CC
         protein is: 5.2, its MW is: 50 kDa.
     NON TER
FT
                   8
                8 AA; 817 MW; A35DD878676B05B1 CRC64;
     SEQUENCE
SQ
  Query Match
                          18.0%; Score 16; DB 1; Length 8;
  Best Local Similarity
                          75.0%; Pred. No. 1.4e+05;
                                0; Mismatches
  Matches
             3; Conservative
                                                1; Indels
                                                                 0; Gaps
                                                                              0;
Qу
           13 EPOG 16
              11 - 1
Db
            3 EPGG 6
```

```
RESULT 43
TKN PHYBI
ID
     TKN PHYBI
                    STANDARD;
                                   PRT;
                                            10 AA.
AC
     P08610;
DT
     01-AUG-1988 (Rel. 08, Created)
DT
     01-FEB-1994 (Rel. 28, Last sequence update)
DΤ
     10-OCT-2003 (Rel. 42, Last annotation update)
DE
     Phyllomedusin.
     Phyllomedusa bicolor (Two-colored leaf frog).
os
OC
     Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC
     Amphibia; Batrachia; Anura; Neobatrachia; Hyloidea; Hylidae;
OC
     Phyllomedusinae; Phyllomedusa.
OX
     NCBI TaxID=8393;
RN
     [1]
RP
     SEOUENCE.
RC
    TISSUE=Skin secretion;
RX
    MEDLINE=70267748; PubMed=5452018;
     Anastasi A., Erspamer G.F.;
RA
RT
     "Occurrence of phyllomedusin, a physalaemin-like decapeptide, in the
RT
     skin of Phyllomedusa bicolor.";
RL
     Experientia 26:866-867(1970).
     -!- FUNCTION: Tachykinins are active peptides which excite neurons,
CC
CC
         evoke behavioral responses, are potent vasodilators and
CC
         secretagogues, and contract (directly or indirectly) many smooth
CC
        muscles.
CC
     -!- SUBCELLULAR LOCATION: Secreted.
     -!- TISSUE SPECIFICITY: Skin.
CC
CC
    -!- SIMILARITY: Belongs to the tachykinin family.
DR
    PIR; S07202; S07202.
    InterPro; IPR002040; Tachy Neurokinin.
DR
DR
    PROSITE; PS00267; TACHYKININ; 1.
KW
     Amphibian defense peptide; Tachykinin; Neuropeptide; Amidation;
KW
     Pyrrolidone carboxylic acid.
FT
    MOD RES
                   1
                          1
                                  PYRROLIDONE CARBOXYLIC ACID.
FT
    MOD RES
                  10
                         10
                                  AMIDATION.
     SEQUENCE
                10 AA; 1189 MW; 3A257D7059D40457 CRC64;
SO
 Query Match
                          18.0%;
                                  Score 16; DB 1; Length 10;
 Best Local Similarity
                          30.0%; Pred. No. 7.4e+03;
 Matches
                                 2; Mismatches
             3; Conservative
                                                   5; Indels
                                                                  0; Gaps
                                                                               0;
            2 QTPQPLLQVM 11
Qу
              | | ::|
Db
            1 QNPNRFIGLM 10
RESULT 44
UPA2 HUMAN
     UPA2 HUMAN
                    STANDARD;
                                   PRT;
ID
                                            10 AA.
AC
     P30088;
DT
     01-APR-1993 (Rel. 25, Created)
DT
     01-APR-1993 (Rel. 25, Last sequence update)
DT
     15-MAR-2004 (Rel. 43, Last annotation update)
DE
     Unknown protein from 2D-page of plasma (Spot 10) (Fragment).
OS
     Homo sapiens (Human).
```

```
OC
     Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC
    Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
OX
     NCBI TaxID=9606;
RN
     [1]
RP
     SEQUENCE.
RC
     TISSUE=Plasma;
RX
    MEDLINE=93092937; PubMed=1459097;
RA
    Hughes G.J., Frutiger S., Paquet N., Ravier F., Pasquali C.,
RA
     Sanchez J.-C., James R., Tissot J.-D., Bjellqvist B.,
RA
     Hochstrasser D.F.;
RT
     "Plasma protein map: an update by microsequencing.";
     Electrophoresis 13:707-714(1992).
RL
CC
     -!- MISCELLANEOUS: On the 2D-gel the determined pI of this unknown
         protein is: 4.4, its MW is: 49 kDa.
CC
     SWISS-2DPAGE; P30088; HUMAN.
DR
FT
    NON TER
                   1
                          1
    UNSURE
FT
                   6
                          6
FT
    NON TER
                  10
                         10
               10 AA; 1079 MW; 51AC54AAB77775B7 CRC64;
SO
     SEQUENCE
 Query Match
                          18.0%; Score 16; DB 1; Length 10;
                          37.5%; Pred. No. 7.4e+03;
  Best Local Similarity
 Matches
             3; Conservative
                                1; Mismatches
                                                  4; Indels
                                                                  0; Gaps
                                                                              0;
           10 VMMEPQGD 17
Qy
              I:I
Db
            1 VXLSPPDD 8
RESULT 45
BPP3 BOTIN
     BPP3 BOTIN
                                   PRT:
ID
                    STANDARD;
                                           11 AA.
     P30423:
AC
DT
     01-APR-1993 (Rel. 25, Created)
     01-FEB-1994 (Rel. 28, Last sequence update)
DT
     28-FEB-2003 (Rel. 41, Last annotation update)
DT
DE
    Bradykinin-potentiating peptide S4,3,2 (10C) (Angiotensin-converting
DE
     enzyme inhibitor).
OS
     Bothrops insularis (Island jararaca) (Queimada jararaca).
OC
     Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC
    Lepidosauria; Squamata; Scleroglossa; Serpentes; Colubroidea;
    Viperidae; Crotalinae; Bothrops.
OC
OX
    NCBI_TaxID=8723;
RN
     [1]
ŔР
     SEQUENCE.
RC
    TISSUE=Venom;
RX
    MEDLINE=90351557; PubMed=2386615;
RA
     Cintra A.C.O., Vieira C.A., Giglio J.R.;
     "Primary structure and biological activity of bradykinin potentiating
RT
RT
     peptides from Bothrops insularis snake venom.";
     J. Protein Chem. 9:221-227(1990).
RL
CC
     -!- FUNCTION: This peptide both inhibits the activity of the
CC
         angiotensin-converting enzyme and enhances the action of
CC
         bradykinin by inhibiting the kinases that inactivate it.
CC
         It acts as an indirect hypotensive agent.
DR
     PIR; C37196; C37196.
KW
     Hypotensive agent; Pyrrolidone carboxylic acid.
```

```
FT
    MOD RES
                                  PYRROLIDONE CARBOXYLIC ACID.
                  1
                         1
               11 AA; 1199 MW; 20B25813C7741777 CRC64;
SQ
     SEQUENCE
                          18.0%; Score 16; DB 1; Length 11;
  Query Match
  Best Local Similarity 50.0%; Pred. No. 8.1e+03;
             3; Conservative
                              1; Mismatches 2; Indels
                                                                 0; Gaps
                                                                             0;
            1 LQTPQP 6
Qу
             1 1:1
            2 LGPPRP 7
Db
RESULT 46
NUDM CANFA
ID
    NUDM CANFA
                    STANDARD;
                                   PRT;
                                           12 AA.
     P54713;
AC
DT
     01-OCT-1996 (Rel. 34, Created)
DΤ
     01-OCT-1996 (Rel. 34, Last sequence update)
DT
     10-OCT-2003 (Rel. 42, Last annotation update)
     NADH-ubiquinone oxidoreductase 42 kDa subunit (EC 1.6.5.3)
DE
    (EC 1.6.99.3) (Complex I-42KD) (CI-42KD) (Fragment).
GN
    NDUFA10.
OS
    Canis familiaris (Dog).
     Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC
OC
    Mammalia; Eutheria; Carnivora; Fissipedia; Canidae; Canis.
OX
    NCBI TaxID=9615;
RN
     [1]
RP
     SEQUENCE.
    TISSUE=Heart;
RC
RX
    MEDLINE=98163340; PubMed=9504812;
RA
     Dunn M.J., Corbett J.M., Wheeler C.H.;
     "HSC-2DPAGE and the two-dimensional gel electrophoresis database of
RT
RT
     dog heart proteins.";
     Electrophoresis 18:2795-2802(1997).
RL
     -!- FUNCTION: Transfer of electrons from NADH to the respiratory
CC
CC
         chain. The immediate electron acceptor for the enzyme is believed
CC
         to be ubiquinone.
     -!- CATALYTIC ACTIVITY: NADH + ubiquinone = NAD(+) + ubiquinol.
CC
CC
    -!- CATALYTIC ACTIVITY: NADH + acceptor = NAD(+) + reduced acceptor.
CC
     -!- COFACTOR: Binds 1 FAD per subunit.
CC
     -!- SUBUNIT: Mammalian complex I is composed of 45 different subunits.
CC
         This a component of the hydrophobic protein fraction.
CC
     -!- SUBCELLULAR LOCATION: Mitochondrial matrix.
    HSC-2DPAGE; P54713; DOG.
DR
     Oxidoreductase; NAD; Ubiquinone; Flavoprotein; FAD; Mitochondrion.
KW
FT
     NON TER
                 12
                         12
SQ
     SEQUENCE
               12 AA; 1284 MW; 3CCD4E2B36EDD737 CRC64;
                          18.0%; Score 16; DB 1; Length 12;
  Query Match
  Best Local Similarity 37.5%; Pred. No. 8.9e+03;
             3; Conservative
                                2; Mismatches 3; Indels
  Matches
                                                                 0; Gaps
                                                                             0;
            6 PLLOVMME 13
Qy
             11 :: 1
```

Db

5 PLAXILGE 12

```
RESULT 47
TM2A METMA
    TM2A METMA
                    STANDARD;
                                   PRT;
                                           12 AA.
AC
     P80652;
     01-OCT-1996 (Rel. 34, Created)
DT
     01-OCT-1996 (Rel. 34, Last sequence update)
DT
     16-OCT-2001 (Rel. 40, Last annotation update)
DT
DΕ
     Alternative tetrahydromethanopterin S-methyltransferase 28 kDa subunit
     (EC 2.1.1.86) (N5-methyltetrahydromethanopterin--coenzyme M
DE
    methyltransferase 28 kDa subunit) (Fragment).
DE
    Methanosarcina mazei (Methanosarcina frisia).
OS
    Archaea; Euryarchaeota; Methanomicrobia; Methanosarcinales;
OC
OC
    Methanosarcinaceae; Methanosarcina.
OX
    NCBI TaxID=2209;
RN
    [1]
RP
     SEQUENCE.
RC
     STRAIN=Goe1 / Go1 / ATCC BAA-199 / DSM 3647 / OCM 88;
RX
    MEDLINE=96370840; PubMed=8774736;
     Lienard T., Becher B., Marschall M., Bowien S., Gottschalk G.;
RA
RT
     "Sodium ion translocation by N5-methyltetrahydromethanopterin:
RT
     coenzyme M methyltransferase from Methanosarcina mazei Gol
RT
     reconstituted in ether lipid liposomes.";
RL
     Eur. J. Biochem. 239:857-864(1996).
CC
     -!- FUNCTION: THIS ENZYME COMPLEX CATALYZES AN INTERMEDIATE STEP IN
CC
         METHANOGENESIS, THE FORMATION OF METHYL-COENZYME M AND
CC
         TETRAHYDROMETHANOPTERIN FROM COENZYME M AND N5-METHYL-
CC
         TETRAHYDROMETHANOPTERIN.
CC
     -!- CATALYTIC ACTIVITY: 5-methyl-5,6,7,8-tetrahydromethanopterin + 2-
CC
         mercaptoethanesulfonate = 5,6,7,8-tetrahydromethanopterin + 2-
CC
         (methylthio) ethanesulfonate.
    -!- SUBUNIT: COMPOSED OF SIX DIFFERENT SUBUNITS.
CC
CC
     -!- SUBCELLULAR LOCATION: Integral membrane protein.
KW
     Transferase; Methyltransferase; Transmembrane; Methanogenesis.
FT
     NON TER
                  12
                         12
     SEQUENCE
                12 AA; 1321 MW; 6DE4A5766232D76B CRC64;
SO
  Query Match
                          18.0%; Score 16; DB 1; Length 12;
  Best Local Similarity
                          25.0%; Pred. No. 8.9e+03;
  Matches
             2; Conservative
                                 5; Mismatches
                                                  1; Indels
                                                                  0; Gaps
                                                                              0;
Qу
            5 QPLLQVMM 12
              :1:1 :::
Db
            5 EPVLPLIV 12
RESULT 48
CRBL ICASP
ID
     CRBL ICASP
                    STANDARD;
                                   PRT;
                                           13 AA.
AC
     P17237;
     01-AUG-1990 (Rel. 15, Created)
DT
     01-AUG-1990 (Rel. 15, Last sequence update)
DT
     28-FEB-2003 (Rel. 41, Last annotation update)
DE
    Chemotactic peptide (I-CP).
os
     Icaria sp. (Ropalidian wasp).
     Eukaryota; Metazoa; Arthropoda; Hexapoda; Insecta; Pterygota;
OC
OC.
     Neoptera; Endopterygota; Hymenoptera; Apocrita; Aculeata; Vespoidea;
OC
     Vespidae; Polistinae; Icaria.
```

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OX
    NCBI TaxID=7495;
RN
     SEQUENCE.
RP
     TISSUE=Venom;
RC
     Yasuhara T., Itokawa H., Suzuki N., Nakamura H., Nakajima T.;
RA
RL
     (In) Izumiya N. (eds.);
     Peptide chemistry 1984, pp.177-182, Protein Research Foundation,
RL
    Osaka (1985).
RL
CC
     -!- FUNCTION: Mast cell degranulating peptide. Induces the chemotaxis
         of neutrophils.
CC
ΚW
    Mast cell degranulation; Chemotaxis; Amidation.
FT
    MOD RES
                  13
                         13
                                  AMIDATION.
     SEQUENCE
                13 AA; 1353 MW; 348DBC7AA30A3768 CRC64;
SQ
  Query Match
                          18.0%; Score 16; DB 1; Length 13;
  Best Local Similarity
                          50.0%; Pred. No. 9.6e+03;
                               2; Mismatches
 Matches
             3; Conservative
                                                 1; Indels
                                                                  0; Gaps
                                                                              0;
            6 PLLQVM 11
Qу
              | | | | ::
            7 PLLGLL 12
Db
RESULT 49
UN37 CLOPA
    UN37 CLOPA
                    STANDARD;
                                   PRT;
                                           14 AA.
AC
     P81358;
DT
     15-JUL-1998 (Rel. 36, Created)
     15-JUL-1998 (Rel. 36, Last sequence update)
DT
DT
     15-JUL-1998 (Rel. 36, Last annotation update)
DE
     Unknown protein CP 37 from 2D-page (Fragment).
OS
     Clostridium pasteurianum.
OC
     Bacteria; Firmicutes; Clostridia; Clostridiales; Clostridiaceae;
OC
     Clostridium.
OX
    NCBI TaxID=1501;
RN
    [1]
RP
     SEQUENCE.
RC
     STRAIN=W5;
RX
    MEDLINE=98291870; PubMed=9629918;
     Flengsrud R., Skjeldal L.;
RA
RT
     "Two-dimensional gel electrophoresis separation and N-terminal
RT
     sequence analysis of proteins from Clostridium pasteurianum W5.";
RL
     Electrophoresis 19:802-806(1998).
     -!- MISCELLANEOUS: ON THE 2D-GEL THE DETERMINED PI OF THIS UNKNOWN
CC
CC
         PROTEIN IS: 5.8, ITS MW IS: 44.7 kDa.
FT
     NON TER
                  14
                         14
     SEQUENCE
                14 AA; 1579 MW; 05384662DEF89210 CRC64;
SQ
  Query Match
                          18.0%; Score 16; DB 1; Length 14;
                          60.0%; Pred. No. 1e+04;
  Best Local Similarity
                                                   1; Indels
  Matches
             3; Conservative
                                 1; Mismatches
                                                                  0; Gaps
                                                                              0;
Qу
           12 MEPOG 16
              :1 11
Db
            7 IEDQG 11
```

```
RESULT 50
COXI THUOB
     COXI THUOB
                    STANDARD;
                                   PRT;
                                           15 AA.
     P80978;
AC
     01-NOV-1997 (Rel. 35, Created)
DT
     01-NOV-1997 (Rel. 35, Last sequence update)
DΤ
     01-NOV-1997 (Rel. 35, Last annotation update)
DT
     Cytochrome c oxidase polypeptide VIc-2 (EC 1.9.3.1) (Fragments).
DE
OS
     Thunnus obesus (Bigeye tuna).
     Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC
     Actinopterygii; Neopterygii; Teleostei; Euteleostei; Neoteleostei;
OC
OC
     Acanthomorpha; Acanthopterygii; Percomorpha; Perciformes; Scombroidei;
OC
     Scombridae; Thunnus.
    NCBI TaxID=8241;
OX
RN
     [1]
RP .
    SEQUENCE.
RC
     TISSUE=Heart;
    MEDLINE=97454291; PubMed=9310366;
RX
RA
     Arnold S., Lee I., Kim M., Song E., Linder D., Lottspeich F.,
     Kadenbach B.;
RA
RT
     "The subunit structure of cytochrome-c oxidase from tuna heart and
    liver.";
RT
RL
     Eur. J. Biochem. 248:99-103(1997).
     -!- FUNCTION: This protein is one of the nuclear-coded polypeptide
CC
CC
         chains of cytochrome c oxidase, the terminal oxidase in
CC
         mitochondrial electron transport.
CC
     -!- CATALYTIC ACTIVITY: 4 ferrocytochrome c + O(2) = 4 ferricytochrome
CC
         c + 2 H(2)0.
    -!- SUBCELLULAR LOCATION: Mitochondrial inner membrane.
CC
    PIR; S77987; S77987.
DR
KW
     Oxidoreductase; Inner membrane; Mitochondrion.
FΤ
    NON TER
                   1
                          1
    NON CONS
                   8
                          9
FT
     NON TER
FT
                  15
                         15
SO
     SEQUENCE
                15 AA; 1696 MW; 4C4C966C73A40294 CRC64;
                          18.0%; Score 16; DB 1; Length 15;
  Query Match
  Best Local Similarity
                          37.5%; Pred. No. 1.1e+04;
             3; Conservative
                                 1; Mismatches
                                                  4; Indels
                                                                  0; Gaps
                                                                              0;
           10 VMMEPQGD 17
Qу
              1 :1
Db
            6 VAKKPMSD 13
```

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Job time : 7.83582 secs